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Coping with Uncertainty

Robust Solutions

 Springer

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Preface

The aim of the series of workshops on “Coping with Uncertainty” (*CwU*) organized at IIASA, Laxenburg, Austria, has been to provide researchers and practitioners from different areas with an interdisciplinary forum for discussing various ways of dealing with uncertainties in diverse areas, including environmental and social sciences, economics, policy-making, management, and engineering. The workshops proved to be successful, especially in cross-disciplinary sharing methods, ideas, and open problems.

Science-based support for addressing the on-going global changes needs solutions for fundamentally new scientific problems, which in turn require new concepts and tools. A key issue concerns a vast variety of practically irreducible uncertainties, including potential extreme events of high multidimensional consequences, which challenge traditional models, and thus require new concepts and analytical tools. This type of uncertainty critically dominates, e.g., the climate change debates. In short, the dilemma is concerned with enormous costs versus massive uncertainties of potentially extreme impacts. Traditional scientific approaches usually rely on real observations and experiments. Yet no sufficient observations exist for new problems, and “pure” experiments, and learning by doing may be very expensive, dangerous, or simply impossible. In addition, the available historical observations are often contaminated by “experimentator”, i.e., past actions, and policies. The complexity of new problems does not allow us to achieve enough certainty just by increasing the resolution of models or by bringing in more links. They require explicit treatment of uncertainties using “synthetic” information composed of available “hard” data from historical observations, the results of possible experiments, and scientific facts as well as “soft” data from experts’ opinions, scenarios, stakeholders, and public opinion. As a result of all these factors, our assessment will always have poor estimates. Therefore, the role of science-based support for addressing the new problems increasingly changes from the traditional “deterministic predictions” analysis to the design of strategies that are robust against the involved uncertainties and risks.

This volume contains contributions based on selected presentation at the *CwU*2007 workshop. The workshop aimed at contributing to a better understanding between practitioners dealing with the safety of complex systems under uncertainty, and scientists working on either corresponding modeling approaches, or on methods that can be adapted for improving the understanding and management of

uncertainty. The focus of the *CwU* 2007 was on novel approaches to supporting robust decision-making and design, especially when uncertainty is irreducible, consequences might be enormous, and the decision process involves stakeholders with diverse interests. Presentations dealt with open problems in this field, limitations of known approaches, novel methods and techniques, or lessons from applications of various approaches. In particular, contributions on the following issues were presented:

- Modeling different types of uncertainty (probabilistic and non-probabilistic)
- The formulation of appropriate deterministic substitute problems for different types of uncertainty
- Robustness of efficient solutions with respect to inherent uncertainties
- Simulation tools (for optimal decision/design under uncertainty)
- Safety and security of humans, environment, and vital infrastructure facing catastrophe risks
- Lessons that can be learned from designing and operating highly reliable systems
- Downscaling and discounting methods for handling spatial and temporal scales
- Benefits and costs of (partial) postponing decisions (aimed at reducing uncertainties)
- Open problems in the adequate treatment of uncertainties
- Concrete applications in economics, finance, engineering, energy, population, air quality, climate change, ecology, forestry, and other environmental problems

The workshop was organized at IIASA in December 2007, jointly by:

- * IIASA – International Institute for Applied Systems Analysis, Laxenburg, Austria
- * Federal Armed Forces University Munich, Germany

The scientific Program Committee included: Yuri Ermoliev, IIASA, Laxenburg (A); Leen Hordijk, IIASA, Laxenburg (A); Marek Makowski, IIASA, Laxenburg (A); Kurt Marti, Federal Armed Forces University Munich (D); Gerhard I. Schuëller, University of Innsbruck (A).

The organizers gratefully acknowledge the support of:

- GAMM – International Association of Applied Mathematics and Mechanics, and
- IIASA – International Institute for Applied Systems Analysis

Their generous support enabled the participation of many researchers who otherwise could not have attended the Workshop.

This volume contains chapters based on selected presentations at the *CwU* 2009 and an introductory short summary of the key issues related to the robust solutions. The chapters are organized into the following four parts:

1. *Modeling of uncertainty* discusses descriptions of uncertainties of different types (probabilities, theory of evidence and possibility, imprecise probabilities, fuzzy sets and variables).

2. *Robust solutions under uncertainty* presents new approaches to discounting applied to evaluation of investments for catastrophic risk management, and to cost-effective and environmentally-safe emission trading under uncertainties, as well as modern quantitative modeling methodologies for analysis of network risks and design of robust networks under uncertainty.
3. *Analysis and optimization of technical systems and structures under uncertainty* deals with state estimation of dynamical systems in case of uncertainties of initial conditions and dynamic parameters described by means of certain ellipsoids, and with the derivation of stochastic linear programs for the reliability-based optimization of plane frames under stochastic uncertainty with respect to external loadings and material parameters.
4. *Analysis and optimization of economic and engineering systems under uncertainty* discusses the variability of the atmospheric deposition of nitrogen in the sea, the treatment of risks and uncertainties in planning agricultural production allocation and expansion, the uncertainty in greenhouse gas emission estimates, consequences of the weather forecasts for the optimal control of agricultural production, and the estimation error in retrieving carbon dioxide column abundances obtained from the GOSAT satellite.

We express our gratitude to all referees, and we thank all authors for the timely delivery of the final version of their contributions to this volume. Furthermore, we thank Ms Elisabeth Löbl of the Federal Armed Forces University Munich for her support in the preparation of this volume. Finally, we thank Springer-Verlag for including the Proceedings in the Springer Lecture Notes Series “LNEMS”.

Munich, Laxenburg
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Kurt Marti
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Contents

- 1 General Remarks on Robust Solutions** 1
Y. Ermoliev, M. Makowski, and K. Marti
References 7
- Part I Modeling of Uncertainty and Probabilistic Issues**
- 2 On Joint Modelling of Random Uncertainty and Fuzzy Imprecision** 11
Olgierd Hryniewicz
 - 2.1 Introduction 11
 - 2.2 Generalizations of Classical Probability and Their Applications in Decision Making 13
 - 2.2.1 Measures of Uncertainty and Criteria of Their Evaluation 13
 - 2.2.2 Probability 15
 - 2.2.3 Dempster–Shafer Theory of Evidence and Possibility Theory 17
 - 2.2.4 Imprecise Probabilities and Their Generalizations 20
 - 2.3 Fuzzy Random Variables and Fuzzy Statistics 22
 - 2.4 Applications of Fuzzy Statistics in Systems Analysis 29
 - 2.4.1 Example 1: Verification of the Kyoto Protocol 29
 - 2.4.2 Example 2: Sequential Testing of a Hypothesis About the Mean Value in the Normal Distribution 31
 - 2.5 Conclusions 33
 - References 35
- 3 On the Approximation of a Discrete Multivariate Probability Distribution Using the New Concept of t -Cherry Junction Tree** 39
Edith Kovács and Tamás Szántai
 - 3.1 Introduction 39
 - 3.2 Preliminaries 40
 - 3.2.1 Notations 40

- 3.2.2 Cherry Tree and t -Cherry Tree 41
- 3.2.3 Junction Tree 42
- 3.3 t -Cherry-Junction Tree 43
 - 3.3.1 Construction of a t -Cherry-Junction Tree 43
 - 3.3.2 The Approximation of the Joint Distribution Over X by the Distribution Associated to a t -Cherry-Junction Tree 44
 - 3.3.3 The Relation Between the Approximations Associated to the First-Order Dependence Tree and t -Cherry-Junction Tree 47
- 3.4 Some Practical Results of Our Approximation and Discussions 50
- References 56

Part II Robust Solutions Under Uncertainty

- 4 Induced Discounting and Risk Management 59**
 T. Ermolieva, Y. Ermoliev, G. Fischer, and M. Makowski
 - 4.1 Introduction 59
 - 4.2 Standard and Stopping Time Induced Discounting 62
 - 4.3 Time Declining Discount Rates 65
 - 4.4 Endogenous Discounting 68
 - 4.5 Dynamic Risk Profiles and CVaR Risk Measure 71
 - 4.6 Intertemporal Inconsistency 73
 - 4.7 Concluding Remarks 75
 - References 76
- 5 Cost Effective and Environmentally Safe Emission Trading Under Uncertainty 79**
 T. Ermolieva, Y. Ermoliev, G. Fischer, M. Jonas, and M. Makowski
 - 5.1 Introduction 79
 - 5.2 Uncertainties and Trends of Carbon Fluxes 82
 - 5.3 Detectability of Emission Changes 84
 - 5.4 Trade Equilibrium Under Uncertainty 86
 - 5.5 Dynamic Bilateral Trading Processes 90
 - 5.6 Computerized Multi-agent Decentralized Trading System 92
 - 5.7 Myopic Market Processes 93
 - 5.8 Concluding Remarks 96
 - References 97
- 6 Robust Design of Networks Under Risks 101**
 Y. Ermoliev, A. Gaivoronski, and M. Makowski
 - 6.1 Introduction 101
 - 6.2 Cooperative Provision of Advanced Mobile Data Services 104
 - 6.3 Simplified Model of the Service Portfolio 106
 - 6.3.1 Description of Services 106

- 6.3.2 Profit Model of an Actor108
- 6.3.3 Service Portfolio: Financial Perspective110
- 6.4 Modeling of Collaborative Service Provision113
 - 6.4.1 Service Provision Capacities114
 - 6.4.2 Risk/Return Industrial Expectations115
 - 6.4.3 Pricing116
 - 6.4.4 Revenue Sharing Schemes.....116
- 6.5 Properties of the Models and Implementation Issues118
- 6.6 Case Study119
- 6.7 Dynamics of Attitudes.....122
 - 6.7.1 Simplified Model: Direct and Indirect Interdependencies123
 - 6.7.2 Model Formulation125
 - 6.7.3 Bayesian Networks and Markov Fields130
 - 6.7.4 Sensitivity Analysis131
 - 6.7.5 General Interdependencies133
- 6.8 Conclusion136
- References136

Part III Analysis and Optimization of Technical Systems and Structures Under Uncertainty

- 7 Optimal Ellipsoidal Estimates of Uncertain Systems: An Overview and New Results141**
 - F.L. Chernousko
 - 7.1 Introduction141
 - 7.2 Reachable Sets142
 - 7.3 Ellipsoidal Bounds145
 - 7.4 Optimality146
 - 7.5 Equations of Ellipsoids148
 - 7.6 Transformation of the Equations.....150
 - 7.7 Properties of Optimal Ellipsoids152
 - 7.8 Generalizations153
 - 7.9 Applications.....154
 - 7.9.1 Two-Sided Estimates in Optimal Control154
 - 7.9.2 Two-Sided Bounds on Time for the Time-Optimal Problem155
 - 7.9.3 Suboptimal Control155
 - 7.9.4 Differential Games156
 - 7.9.5 Control of Uncertain Systems157
 - 7.9.6 Other Applications157
 - 7.9.7 State Estimation in the Presence of Observation Errors....158
 - 7.10 Ellipsoidal vs. Interval Analysis159
 - 7.11 Conclusions160
 - References160

8	Expected Total Cost Minimum Design of Plane Frames by Means of Stochastic Linear Programming Methods	163
	Kurt Marti	
8.1	Introduction	164
8.1.1	Plastic Analysis of Structures	164
8.1.2	Limit (Collapse) Load Analysis of Structures as a Linear Programming Problem	165
8.1.3	Plastic and Elastic Design of Structures	167
8.2	Plane Frames	168
8.2.1	Yield Condition in Case of $M - N$ -Interaction	173
8.2.2	Approximation of the Yield Condition by Using Reference Capacities	180
8.3	Stochastic Optimization	183
8.3.1	Violation of the Yield Condition	184
8.3.2	Cost Function	185
8.3.3	Choice of the Cost Factors	186
8.3.4	Total Costs	187
8.3.5	Discretization Methods	189
8.3.6	Complete Recourse	190
	References	191

Part IV Analysis and Optimization of Economic and Engineering Systems Under Uncertainty

9	Uncertainty in the Future Nitrogen Load to the Baltic Sea Due to Uncertain Meteorological Conditions	195
	Jerzy Bartnicki	
9.1	Introduction	195
9.2	Nitrogen Emissions	198
9.2.1	National Emission Ceilings According to EU NEC Directive	198
9.2.2	National Emission Ceilings According to Gothenburg Protocol	199
9.2.3	Nitrogen Emission Projections Used in the Model Runs	200
9.3	Computed Nitrogen Depositions for 2010	201
9.3.1	Unified EMEP Model	202
9.3.2	Calculated Depositions to Sub-basins and Catchments of the Baltic Sea	203
9.4	Uncertainty Due to Meteorological Variability	203
9.5	Conclusions	207
	References	207

10 Planning Sustainable Agricultural Development Under Risks	209
G. Fischer, T. Ermolieva, and L. Sun	
10.1 Introduction	209
10.2 Cooperation and Co-existence for Risk Sharing	211
10.3 Agricultural Planning Under Risks	214
10.3.1 A Simulation Model	214
10.3.2 A Simplified Production Model	216
10.3.3 A Rebalancing Production–Allocation Algorithm	217
10.4 Stochastic Production Allocation Model	218
10.5 Numerical Experiments	221
10.6 Conclusions	225
References	226
11 Dealing with Uncertainty in GHG Inventories: How to Go About It?	229
Matthias Jonas, Thomas White, Gregg Marland, Daniel Lieberman, Zbigniew Nahorski, and Sten Nilsson	
11.1 Introduction	230
11.2 Does Uncertainty Matter?	232
11.3 State of the Art of Analyzing Uncertain Emission Changes	233
11.4 How to Deal with Uncertainty?	238
11.5 Conclusions	241
References	242
12 Uncertainty Analysis of Weather Controlled Systems	247
K.J. Keesman and T. Doeswijk	
12.1 Introduction	247
12.2 Preliminaries	249
12.2.1 Bulk Storage Model	249
12.2.2 Weather Forecasts	249
12.2.3 Cost Function	250
12.2.4 Receding Horizon Optimal Control	250
12.3 Weather Forecast Uncertainty and Error Analysis	252
12.3.1 Open Loop Evaluation	252
12.3.2 Closed Loop Evaluation	254
12.4 Discussion	256
12.5 Concluding Remarks	257
References	257
13 Estimation of the Error in Carbon Dioxide Column Abundances	259
Mitsuhiro Tomosada, Koji Kanefuji, Yukio Matsumoto, Hiroe Tsubaki, and Tatsuya Yokota	
13.1 Introduction	259

13.2	Trace Gas Measurement by Satellite Remote Sensing.....	261
13.2.1	Observations of Trace Gases with Various Sensors	261
13.2.2	GOSAT Mission	262
13.2.3	Previous Error Analysis	264
13.3	Error Evaluation and Results.....	266
13.3.1	Retrieval Method.....	266
13.3.2	Error Evaluation	268
13.3.3	Error Evaluation Results.....	270
13.4	Conclusions	276
	References.....	276

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Chapter 1

General Remarks on Robust Solutions

Y. Ermoliev, M. Makowski, and K. Marti

We summarize here the background and key concepts related to robust solutions in the context of supporting decision-making for problems characterized by deep uncertainties, which also were in the focus of the previous workshops on *Coping with uncertainty*, see, e.g., [3]. Although such problems are fundamentally different from statistical decision models, yet basic ideas of robust statistics are applicable to methods supporting robust decision-making under uncertainty. The main new issues are concerned with a proper representation of uncertainty, and its interactions with decisions. In particular, a key issue is the sensitivity of robust decisions with respect to low probability catastrophic events, that are of critical importance for analyzing global change problems. Robust decisions for problems exposed to extreme catastrophic events are essentially different from over-simplified decisions that ignore such events. Specifically, a proper treatment of extreme/rare events requires new paradigms of rational decisions, new performance indicators, and new spatio-temporal dimensions of heterogeneous interdependencies including network externalities and risks. This, in particular, needs new approaches to downscaling, upscaling and discounting.

Global change processes, in particular climate change, involve inherently unpredictable complex interactions between natural and human-created systems therefore proper modeling of these processes must rely on adequate treatment of uncertainties, and their effects on human's decisions. Traditional natural science models are based on relations whose validity is estimated from repetitive experiments and observations. If experiments do not affect the underlying relations, then repetitive observations allow to derive them by using the statistical decision theory. Unfortunately, human-created processes do not follow fixed relations. Elements of these processes change their dimensions and structure. For example, introduction of new

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technologies may increase or reduce uncertainties, risks, critical thresholds and discontinuities. Exact identification of global climate change processes is impossible because such processes are non-stationary, have delayed responses, and human or natural actions may have catastrophic irreversible consequences.

Under inherent uncertainty of heterogeneous processes the role of integrated models rests on the ability to guide comparative analysis of rational decisions. Although exact evaluations are impossible, the preference structure among decisions can be a stable basis for a relative ranking of alternatives in order to design robust policies, which must be in a sense optimal against all relevant uncertainties. It is commonly known that finding out (without exact measurement) which of two parcels is the heavier is much easier than evaluating weight of each parcel.

The term “robust” was introduced into statistics in 1953 by Box [2] and acquired recognition after the publication of a path-breaking paper by Huber [5]. As Huber admits, researchers had long been concerned with the sensitivity of standard estimation procedures to “bad” observations (outliers), and the word “robust” was loaded with many, sometimes inconsistent connotations, frequently for the simple reason of conferring respectability on it. Appeal for robustness [4] probably dates back to pre-history of statistics. A distant outlier in observations ruins the least square analysis, therefore rejection of outliers is a sort of robust statistical procedure. The discussion about the rejection of outliers is at least as old as the 1777 publications of Daniel Bernoulli [1]. The mean is not robust to outliers, whereas the median is robust. Therefore, switching from the mean to the median for long-tailed data increases robustness. This is also equivalent to switching from quadratic (least square) smooth optimization to non-smooth optimization principles.

According to Huber [5], “...any statistical procedure...should be robust in the sense that small deviations from the model assumptions should impair the performance only slightly”. This concept of robustness corresponds to standard mathematical ideas of continuity and stability: when disturbances become small, the performance of the perturbed model also deviates slightly. In other words, a robust procedure is in a certain sense optimal with respect to all uncertainties from a neighborhood of the model. Huber introduced rigorous notions of robustness based on probabilistic minimax approach and Choquet capacity (imprecise probabilities), which lead to specific non-smooth stochastic optimization models. By using appropriate neighborhoods of probability measures (e.g., with respect to ϵ -contaminated probabilities, Levy distance, or Kolmogorov distance), he derived robust estimators optimizing the worst that can happen over the neighborhood of the model with respect to a certain performance indicator. Neighborhoods of probability measures can also be characterized by Choquet capacities, i.e., functions which define sets of probability measures by taking all probabilities which lie below (or above) a capacity (point-wise).

These basic ideas of robust statistics as well as the infinitesimal robustness introduced by Hampel [4] are also used for more general decision problems under uncertainty. In particular, the infinitesimal approach is based on the fact that many statistics and solutions of general decision models can be considered as functionals in the space of probability measures. The robustness information is then provided

by the inference functions, roughly speaking, a derivative (in the space of probability measures) of a statistic or a performance indicator at an underlying distribution. There are also important concepts of Bayesian and non-Bayesian robustness, where we need not only robustness against deviations from the given parametric model, but also against uncertainties of the prior distributions.

The word “robust” has become fashionable in statistical decision theory and other disciplines dealing with data analysis and decisions [4], in particular, for dealing with questions such as: Which data are of critical importance and should be examined with a special care? What methods provide the greatest safety? How safe are results of a model that is known only approximately?

As the concept of statistical robustness is in a sense similar to the problem of local stability of dynamic systems, the robustness in deterministic control theory was introduced as an additional requirement on the stability of optimal trajectories. In other words, additional constraints were introduced in the form of a stability criterion. Optimization theory provides tools for analyzing and solving various decision making problems. For deterministic models robustness is defined similar to probabilistic minimax robustness in statistics: to optimize the worst that can happen to performance indicators over solutions x that satisfy feasibility constraints for all admissible values of uncertainty $\omega \in \Omega$. The set Ω is often characterized by a finite number of scenarios or simple sets such as intervals or ellipsoidal uncertainty which, in a sense, attempt to substitute for normal probability distributions in a simple but inconsistent with statistical analysis manner. It is clear that this type of deterministic worst-case robustness leads to extremely conservative decisions.

After the word robust become fashionable in statistics, it is being used in many senses, e.g., “quantitative robustness”, “ Π -robustness”, “B-robustness”, see, e.g., [4]. The situation becomes more complex for general decision problems under uncertainty dealing with quite different decision situations which may have a vast variety of different facets of robustness. Therefore, in order to avoid dangerous confusions, the term “robust” must be precisely defined in every specific context. This is similar with other notions whose meaning depends on the context, e.g., fairness, efficiency, optimality. For example, a decision is optimal only with respect to precisely specified conditions.

Statistical decision theory deals with situations in which a model of uncertainty and corresponding optimal solution are defined by a sampling model characterized by a probability measure P with an unknown vector of “true parameters” x^* . Vector x^* defines a desirable optimal solution, its performance can be observed from the sampling model, and the problem is to recover x^* from these data. Potential estimates of x^* define feasible solutions x of a statistical decision problem. It is essential that x does not affect the sampling model P so that the optimality and robustness of solutions can be evaluated by a distance from x^* by using its observable performance.

The general problems of decision making under uncertainty deal with fundamentally different situations. The model of uncertainty, feasible solutions, and performance of the optimal solution are not given and all of these have to be characterized from the context of the decision making situation, e.g., socio-economic,

technological, environmental, and risk considerations. As there is no information on true optimal performance, robustness cannot be also characterized by a distance from an observable true optimal performance. Therefore, the general decision problems may have rather different facets of robustness. In particular, probabilistic minimax solutions may seem too pessimistic or too optimistic for coping with potentially catastrophic events. Therefore, other concepts of robustness are required. Such concepts may be based on, e.g., expected utility maximization, stochastic optimization, stochastic minimax models.

In the presence of uncertainty, any related decision x results in multiple outcomes such as costs, benefits, damages, and risks, as well as indicators of fairness, equity, and environmental impacts. The outcomes depend not only on decisions x but also on uncertainty characterized by $\omega \in \Omega$, where Ω , denotes a set of admissible scenarios.

Scenario analysis is often used as a straightforward approach to find a decision that is “optimal” with respect to all scenarios by attempting to solve the decision problem for all possible scenarios. Unfortunately, a given decision x for different scenarios ω may have rather contradictory outcomes which do not really tell us which decision is reasonably good for all scenarios.

In 1738 mathematician Daniel Bernoulli (see discussion in [7]) introduced the concept of expected utility maximization as a rule for choosing decisions under multiple outcomes. It is assumed that all outcomes $g_i(x, \omega)$, $i = 1, \dots, K$ can be summarized in a single measure of preferability, e.g., a monetary payoff denoted by: $q(x, \omega) = Q(g_1, \dots, g_K)$. The standard expected utility model suggests to choose a decision x that maximizes an expected utility function

$$U(x) = Eu(q(x, \omega)) = \int u(q(x, \omega))P(d\omega),$$

i.e., in a sense, for all $\omega \in \Omega$, where $u(\cdot)$ is a utility associated with an aggregate outcome $q(x, \omega)$. The shape of u defines attitudes to risks. This model presupposes that one can rank the alternative scenarios ω according to weights – objective or subjective probability measure P .

The use of a probability measure as a degree of belief was formalized by Ramsey [6]. Savage published [7] a thorough treatment of expected utility maximization based on subjective probability as a degree of belief. As a result of this work the use of probability measure became a standard approach for modeling uncertainty by using “hard” observations and “soft” public and expert opinions in a consistent way within a single model. Although a decision maximizes the expected utility, in a sense, for all scenarios, still it cannot be considered to be a robust solution. The shortcomings of the expected utility model are well known. Generally, it is practically impossible to find a utility function that enables satisfactory aggregation of various attributes in one preferability measure, including attitudes to different risks, the distributional aspects of gains and losses, the rights of future generations, and responsibilities for environmental protection.

For complex problems it is natural that different performance indicators should be used to evaluate robustness of the integrated system in the same way as we use indicators of health (e.g., temperature and blood pressure for humans). An expected utility model is a specific case of stochastic optimization (STO) models that use various performance indicators $f_i(x, \omega)$, $i = 1, \dots, m$, one of which can be the expected utility (disutility). These indicators depend on outcomes $g_k(x, \omega)$, $k = 1, \dots, K$, on x and ω , i.e.,

$$f_i(x, \omega) := q_i(g_1, \dots, g_K, x, \omega).$$

A rather general STO problem is formulated as optimization (maximization or minimization) of the expectation function

$$F_0(x) = E f_0(x, \omega) = \int f_0(x, \omega) P(d\theta)$$

subject to constraints

$$F_i(x) = E f_i(x, \omega) = \int f_i(x, \omega) P(d\theta) \geq 0, \quad i = 1, \dots, m.$$

The choice of proper indicators $f_i(x, \omega)$ and outcomes $g_k(x, \omega)$, $k = 1, \dots, K$, is essential for the robustness of x . Globally or regionally aggregated outcomes are less uncertain but they may not reveal potentially dramatic heterogeneities induced by global changes on individuals, governments, and the environment. For instance, an aggregate income or growth indicators may not reveal an alarming gap between poor and rich regions, which may cause future instabilities.

By choosing appropriate outcomes $g_k(x, \omega)$ and functions $f_i(x, \omega)$, STO models allowing a natural and flexible way to represent various risks, abrupt changes, spatio-temporal heterogeneities, equity constraints and the sequential resolution of uncertainty in time. Often, under proper robustness requirements, $f_i(x, \omega)$ are analytically intractable, non-smooth, and even discontinuous functions, and probability measure P is chosen from a feasible set, thus is imprecise; moreover, P often depends on x , which is essential for modeling endogenous extreme events and catastrophic risks, and non-Bayesian robustness. It is often practically impossible to identify uniquely subjective (and objective) probability as a degree of beliefs. Most people cannot clearly distinguish between probability ranging roughly from 0.3 to 0.7. Decision analysis often has to rely on imprecise statements, for example, that event e_1 is more probable than event e_2 or that the probability p_1, p_2 of event e_1 or of event e_2 is greater than 50% and less than 90%. Therefore, feasible sets of probabilities may be represented by inequalities such as $p_1 \geq p_2, 0.5 \leq p_1 + p_2 \leq 0.9$. As in robust statistics, the robust solutions of general decision models can be derived by using worst-case (for a given decision) probability distribution from the feasible sets of distributions satisfying constraints of STO model. The applicability of the

infinitesimal approaches is facilitated by the fact that solutions of STO models are functionals of underlying probability measures.

In contrast to the expected utility based approaches, the STO models support integrated solutions composed of anticipative ex-ante and adaptive ex-post decisions, which in turn allows to model flexible decision processes with adaptive adjustments of anticipative decisions when new information becomes available. The ability of STO models to incorporate both anticipative ex-ante and adaptive ex-post decisions induces non-smooth criteria and risk aversion among ex-ante decisions that implicitly depends on input data and practically cannot be characterized by an exogenous utility function. In particular, even in the simplest linear model (see [3] and chapters of Part II of this volume) the co-existence of ex-ante and ex-post decisions induces quantile-type risk measures. Therefore, this approach allows to substitute mean values by median or/and other quantiles without destroying convexity (concavity) of models.

A fundamentally new aspect of robustness for general decision problems is the sensitivity (or discontinuity) of robust solutions with respect to extreme (low probability and high consequences) events affecting large territories and communities, see [3]. Although potential extreme events with large-scale catastrophic impacts dominate the global change policy discussions there is no agreement on the corresponding measures of the changes and the impacts. For example, a commonly used measure is the projected global mean temperature change, which falls within the difference between the average temperature of cities and their surrounding rural areas, and cannot be reliable indicator of climate change. Actually, global climate change impacts can be properly evaluated only in terms of local temperature variability and related extreme events, in particular, heat waves, floods, droughts, wind-storms, diseases, and sea-level rise.

Unfortunately, ignorance of extreme events explicitly or implicitly through inadequate treatments of their impacts can be dramatically misleading. The latter is especially dangerous since it provides an illusion of truly comprehensive analysis. A 500-year disaster, e.g., an extreme flood that occurs on average once in 500-years is often considered as irrelevant for many future generations, but actually it may occur next year and destabilize economy for many years. However, existing extremal value theory deals primarily with independent variables quantifiable by a single number, e.g., money. Therefore, the use of this theory may also be misleading, because catastrophes are definitely not quantifiable events in this sense. They have different patterns, spatial and temporal dimensions and induce heterogeneity of losses and gains, which exclude the use of aggregate space-less characteristics. Globally, an average resident may even benefit from some climate change scenarios, while some regions would be simply wiped out.

Ignorance of spatio-temporal details leads to models and solutions which are insensitive to large-scale extreme events (outliers), therefore formally can be viewed as robust solutions demonstrating insignificance of threats related to on-going global changes, say, climate change. The approaches presented in [3] and in this volume allow to properly address issues of robustness of solutions for problems exposed to catastrophic events. In particular, probabilistic maximin (minimax) robustness

may not be sufficient for this. A more general approach is the combination of the probabilistic and so-called stochastic maximin approach maximizing

$$\min_{p \in P} E \min_{z \in Z} f_0(x, y, z, \xi)$$

under constraints of STO model, where x are decision variables, and scenario ω is defined by the vector of variables y, z , and ξ : $\omega = (y, z, \xi)$. The components of ω are: $y \in Y$, variables representing uncertainty ranked by an objective or subjective probability measure from P ; $z \in Z$, variables representing potential extreme random scenarios, as in the extremal value theory; ξ are variables ranked by a fixed probability measure as in the basic STO models. The concept of stopping time allows to focus the analysis on the least-probable and the most destructive extreme events. As shown in [3], there are strong connections between the stopping time concept and the stochastic maximin.

In the absence of sufficient information, models play a key role in comparative analysis of alternative solutions for designing robust policies. Any policy analysis focuses attention on situations where processes can be changed by decisions that should be selected in the best possible manner. In reality, a proper structure of models, e.g., sets of proper decision are also uncertain and they can be specified through a dialogue of users with models. Specification of such models include use of quantiles, thresholds, and stopping times, which in turn require specific non-smooth stochastic optimization methods. Effective analysis of such models requires development of specific fast adaptive Monte Carlo optimization procedures.

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Part I
Modeling of Uncertainty
and Probabilistic Issues

Chapter 2

On Joint Modelling of Random Uncertainty and Fuzzy Imprecision

Olgierd Hryniewicz

Abstract The paper deals with the problem of the mathematical description of uncertainties of different type. It has been demonstrated by many authors that the theory of probability is not always suitable for the description of uncertainty related to vagueness. We briefly present some of the most promising theories which have been recently proposed for coping with this problem. Then, we concentrate our attention on the application of fuzzy random variables which seem to be very useful for the joint modelling of random uncertainty and fuzzy imprecision, and for the statistical analysis of imprecise data. The application of the statistical methodology for fuzzy data, called fuzzy statistics, is illustrated with two practical examples, typical for the problems of systems analysis. First example is devoted to the problem of the estimation of greenhouse gases inventories. In the second example, typical for the problems of making decisions using small amount of available data, we show how fuzzy approach can be used for the improvement of sequential statistical tests.

2.1 Introduction

Coping with uncertainty is an important problem in many areas of science, but in systems analysis and decision sciences it becomes a really crucial one. In both these branches of science uncertainty is always present, as systems analysts and decision makers have never full information about past, current and future “states of the world”. Thus, their information about consequences of proposed by them actions, which is necessary, e.g. for finding optimal decisions, is rarely precise and fully reliable. Moreover, they usually cannot present descriptions of processes of their interest with accuracy which is typical, e.g. for physics, chemistry or astronomy. Therefore, they need to use a formal language that could be used for sufficiently precise description of uncertain events, actions, etc. For many decades it appeared

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to the majority of scientists that the theory of probability and mathematical statistics is the only methodology that should be used for the formal description of uncertainty. However, during last few decades many scientists working in such areas like psychology, economic sciences, quantum physics, artificial intelligence, etc. have raised questions about possible inadequacy of the classical (Kolmogorov's) theory of probability when applied for solving their particular problems.

This situation is far from being unexpected. If we look at any good dictionary of, e.g. English we can find that the word "probable", which definitely describes uncertainty, has many synonyms and other related words. For example, such words like "possible", "plausible", and expressions like "likely to be true", "hopeful", "to be expected" (and their antonyms) are used for the description of a state on uncertainty. One may expect that they are used for expressing slightly different types of uncertainty, and are not fully exchangeable with the word "probable". The differences between their meanings raised doubts among philosophers and mathematicians about the role of the classical theory of probability as the sole mathematical language for the description of uncertainty.

Basic problems with the applicability of the classical theory of probability inspired mathematicians who proposed other, not necessarily equivalent, theories of probability. Some of them are described in a classical book of Fine [24]. Other doubts were raised by the founder of the theory of fuzzy sets L.A. Zadeh who claimed that the classical theory of probability cannot describe uncertainty related to innate imprecision of many notions and ideas expressed in a plain human language. In his seminal paper [95] Zadeh proposed to use the formalism of fuzzy sets as the formal language of the theory of possibility. Another criticism of the classical probability came from economists and psychologists. The Nobel Prize winner in economics H. Simon in his book [77] noticed that people do not make their decisions according to the principle of expected utility which is based on the classical theory of probability. Other doubts were raised by other Nobel Prize winners Tversky and Kahneman who noticed in their works (see, for example, [83]) that probabilities evaluated by humans are not necessarily additive, as it is assumed in the classical theory of probability. An interesting description of theoretical and practical problems with the applicability of the classical theory of probability can be found in the paper by Bordley [5] who also noticed that this theory is in a certain sense incompatible with the quantum physics.

The existence of many extensions and modifications of the classical theory of probability creates problems for systems analysts and decision makers who are expected to model systems in presence of uncertainties of different types. In the second section of this paper we present a very brief description of several generalizations of classical probability. This has to be done in order to set borders between those areas where classical probability is still the best (and probably the only) mathematical model of uncertainty and the areas where its generalizations are needed. We claim that in the majority of practical cases the combination of classical probability and Zadeh's theory of possibility is sufficient for the description of complex systems and making decisions. We describe this methodology in the third section of

the paper. Examples of the application of fuzzy random models and fuzzy statistics are given in the fourth section of the paper. We show that in case of information that has both random and imprecise nature some additional indices, like possibility and necessity measures, are indispensable for a correct description of decision making problems.

2.2 Generalizations of Classical Probability and Their Applications in Decision Making

2.2.1 Measures of Uncertainty and Criteria of Their Evaluation

Close analysis of the human perception of uncertainty reveals that this concept does not have one, unanimously approved, interpretation. Zimmermann [96] notes that any definition of uncertainty has to be to some extent arbitrary and subjective. He proposes the following one:

Uncertainty implies that in a certain situation a person does not dispose about information which quantitatively and qualitatively is appropriate to describe, prescribe or predict deterministically and numerically a system, its behavior or other characteristics.

This definition is definitely technology-oriented. One may say that everything what prevents us to describe reality in a deterministic way may be considered as a facet of uncertainty. Thus, there exist many different causes of uncertainty. Zimmermann [96] lists the following: lack of information (quantitative or qualitative), abundance of information, conflicting evidence, ambiguity, measurement, and belief. It is not surprising that he does not believe that the general theory of uncertainty which is able to describe these completely different sources of uncertainty exists, and appropriate mathematical models should be context-dependent. They could be formulated either as different generalizations of probability, or may be formulated in another way, such as Pawlak's rough set theory (see, e.g. [67]) or convex modeling proposed by Ben-Haim and Elishakoff [3].

If we look at different theories of uncertainty we can notice that they can be divided into two general groups: those based on methods of mathematical logics (such as the rough sets theory) or those based on the theory of probability and its generalizations. In this paper we restrict our interest only to the second one. This restriction arises from a fact that in the majority of practical cases the classical theory of probability is sufficient for the mathematical description of uncertainty. This popularity of classical probabilistic models of uncertainty makes many specialists to believe that classical probability is the only mathematical theory that is sufficient for the formal description of uncertainty. Insufficiency of this approach was noticed only recently, mainly by specialists in decision-making or expert systems. Peter Walley, who is the one of the most prominent persons representing that group of scientists, presents the list of pertinent mathematical models, in order of their generality [88] (in parentheses, there are given the most important, according to Walley,

references, and indicated by him typical areas of application):

- Possibility measures and necessity measures ([18, 95], vague judgments of uncertainty in natural language)
- Belief functions and plausibility functions ([12, 76], multivalued mappings and non-specific information)
- Choquet capacities of order 2 ([8, 13, 45], some types of statistical neighborhood in robustness studies, and various economic applications)
- Coherent upper and lower probabilities ([45, 56, 78], personal betting rates, and upper and lower bounds for probabilities)
- Coherent upper and lower previsions ([86, 87, 92], buying and selling prices for gambles, upper and lower bounds for expectations, and envelopes of expert opinions)
- Sets of probability measures ([4, 36, 58], partial information about an unknown probability measure, and robust statistical models)
- Sets of desirable gambles ([86, 91, 93], preference judgments in decision making)
- Partial preference orderings ([34, 86], preference judgments in decision making)

Walley [88] also notices:

- Partial comparative probability orderings ([25, 47, 49], qualitative judgements of uncertainty)

In order to evaluate and compare all competing theories of uncertainty (including classical probability) Walley [87] proposes to take into account the following criteria:

- (a) Interpretation
- (b) Imprecision
- (c) Calculus
- (d) Consistency
- (e) Assessment
- (f) Computation

The proposed measure of uncertainty has to be sufficiently easy to understand by its users. For example, conclusions inferred from the application of the theory should be clear enough to be useful for making decisions. It should be able to model partial or complete ignorance, reflected, for example, in imprecision of statements of natural language. There should be rules for merging uncertainties, updating, and using them in inferential processes. There should be methods for the evaluation of coherence of all assessments formulated using the theory and its assumptions. A useful theory of uncertainty should provide guidance how to make assessments about uncertain events and handle imprecise judgments of different types. Finally, it should be computationally feasible. More comprehensive interpretation of the criteria presented above and their practical and theoretical importance can be found in [87].

All these requirements may have different importance in different applications, and none of the existing theories and measures of uncertainty fulfills them sufficiently well. For example, the classical theory of probability does not meet

sufficiently well criteria (b) and (e), and for this reason philosophers, mathematicians, economists, psychologists, and specialists in expert systems have been making a lot of efforts in order to introduce more general, and more useful in specific applications, theories of uncertainty. In the following subsections of this section we present very brief description of some of these theories. This presentation is needed, in our opinion, for the understanding of limits which still exist if we try to cope with uncertainty inherent in the analysis of complex systems.

2.2.2 Probability

The theory of probability is the best known, and the most frequently used in practice, theory of uncertainty. Many specialists claim that it is the only consistent theory that describes all types of uncertainty, both *aleatoric* describing randomness observed in repeated experiments and *epistemic* describing uncertain information provided by human beings (e.g. experts). It has a well established mathematical formulation, and efficient methods for processing information described using probabilistic concepts. However, despite its four hundred years lasting history its fundamentals are still subject to different interpretations and controversies. In general, there exist two different interpretations of the classical probability: an objective “frequentist” interpretation, based on the analysis of empirical observations of series of events, and subjective “Bayesian” approach, based of subjective assessment of probabilities of events. It is interesting that even in the 1960s the second approach was dismissed as “non-scientific” by the majority of statisticians, and not present in nearly all popular textbooks. On the other hand, the supporters of the Bayesian approach presented in books of Savage [74] and de Finetti [26, 27] pointed out apparent incoherences inherent for the frequentist approach (see, for example, an excellent monograph by Lindley [59]).

The basics of the theory of probability are well described in all textbooks on probability and statistics. Therefore, there is no need to present them in details in this paper. However, we are going to point out those assumptions of this theory which are criticized by some authors who see them as main reasons of discrepancies between theory and practice of coping with uncertainties.

According to Kolmogorov’s theory of probability there exists a sample space Ω consisting of *disjoint* elements, called elementary states, or simply states. These states need not be necessarily observable. Then, Kolmogorov postulates a Borel-field set B consisting of some, but not necessarily all, subsets of Ω . The elements of B are called events and represent observable outcomes of actual or hypothetical experiments. Probabilities are assigned only to elements of B , so they are not assigned to those states of Ω that do not belong to B . One can argue, however, that those events which cannot be included in an appropriate σ -algebra are irrelevant and may be treated as Lebesgue null sets. A counter-argument to this point of view may be the following: in presence of partial information (or partial ignorance) we may not be entitled to make such claims. The consequence of these assumptions

is far-reaching, and it does not depend upon the interpretation of probability. It means that every event can be *precisely* described using the elements of Ω . Another consequence refers to the feature which Walley [87] calls “Bayes dogma of precision”. According to this feature of the theory of probability, every uncertain entity (physical object, value of a parameter, etc.) can always be described by a *precisely* defined probability distribution. Some supporters of this assumption even question the reality of situations when the only available information about uncertainty can be presented in form of an interval (see quotations presented in the paper [23] which summarizes discussions on the problems related to the treatment of epistemic uncertainty). When probabilities are assessed by frequencies of observed precisely defined events there are no fundamental problems with the accuracy of their evaluation. However, when they are assessed subjectively (and we have to remember that according to the followers of the Bayesian probability and statistics it is the only *coherent* way of doing this) it is assumed that they may be interpreted as precisely defined fair betting rates. The behavioral interpretation of probabilities in terms of fair betting rates was originally introduced by de Finetti (see [27]) who has shown that betting in favor of an event A against its complement A^C will not lead to sure loss only if betting odds are $P(A)$ to $1 - P(A)$, where $P(A)$ is equal to the probability of event A . Moreover, from the postulate of fair betting rates and some additional coherence requirements one can derive that probability is nonnegative, normalized, and finitely-additive set function. In addition, Savage [74] proved that the theory of subjective probabilities constitutes the basis for an axiomatized and coherent theory of decision-making.

Despite of all these unquestionable advantages, empirical observations show however, that in the presence of partial or full ignorance about events of interest such precise assessments of probability cannot be made. Moreover, the actual behaviour of decision-makers differs from that prescribed by the theory based on classical precise probabilities. Bordley [5] argues that this apparent inconsistency is due to the fact that real decision-makers take into account incompleteness of their information. This leads us (and many other researches working in the area of decision sciences) to the conclusion that in case of imprecisely defined states and events it is not possible to obtain precise values of their probabilities, and hence to make precise prescriptions in decision-making processes. A counterargument to this opinion presented by rather dogmatic followers of the classical Bayesian approach to probability is the following: their theory shall be considered as the normative one, and all the differences between the theory and the actual human behaviour are always due to human weakness and shall be overcome by using more precise measurements and precise problem formulation. In many practical cases this is definitely true. However, even in principle this standpoint can be questioned using the results from quantum physics. Bordley [5] shows that as the consequence of the Heisenberg Uncertainty Principle some events cannot be precisely observed, and in such a case precise probability statements are in principle impossible. Therefore, generalizations of classical probability are necessary if we want to deal with imprecisely defined events and with partial information about probabilities of their occurrence.

It has to be stressed, however, that once the probability distributions describing uncertain phenomena have been defined, the mechanisms of the theory of probability are sufficient for further analysis of uncertain phenomena. For this reason probabilistic methods, such as Bayesian methods, are generally used in practice. Basic principles of the application of the Bayesian methods are described in classical textbooks by Raiffa and Schlaifer [69] and De Groot [11]. Examples of particular applications are presented in numerous publications from virtually all areas of science. The only point of criticism formulated against this approach stems from the fact that according to some scientists the existing imprecise or incomplete information does not allow us to propose those precisely defined probability distributions without making some additional assumptions. Other approaches proposed in order to cope with this problem are presented in the next sections of this paper.

2.2.3 Dempster–Shafer Theory of Evidence and Possibility Theory

The notion of *possibility* attracted attention of philosophers, economists, logicians, etc. Dubois and Prade [20] notice that first attempts to formalize the concept of possibility were made in the late 1940s by the economist Shackle [75] who proposed a calculus of “potential surprise” as the base for decision-making. The works of many authors, who have noticed the deficiency of the theory of probability in dealing with many practical problems have led to more or less independent formalizations of two similar theories of uncertainty: Dempster–Shafer theory of evidence and possibility theory.

The concept of possibility can be interpreted in different ways. For example, it can be understood as an objective notion or as an epistemic and subjective one. Zadeh [95] understands possibility as objective feasibility; an objective measure of physical easiness to achieve a certain goal. By his famous example of a possibilistic statement, “it is possible for Hans to eat six eggs for breakfast”, he shows an exemplary information which is difficult, or even hardly possible, to be formalized using theory of probability. This type of interpretation of possibility is closely related to the idea of preference. Alternatives that are more easily achieved (more feasible) are usually more preferred. This relation has been described in details in the paper by Dubois, Fargier, and Prade [14]. Second interpretation of possibility is an epistemic one, and is given in terms of plausibility. An event is fully plausible when its occurrence does not create any surprise. This type of interpretation has subjectivistic nature and means that possibility may represent consistency of the observed event with available knowledge. Possibility, understood as plausibility of an event, may also have an objectivistic interpretation, and can be evaluated from the observations of upper bounds of frequency of its occurrence [17]. There also exists a deontologic interpretation of possibility (something is possible when it is allowed by law), but satisfactory formal description of this type of possibility has not been proposed yet. All those interpretations, which in the majority of practical cases are hardly

exclusive, lead to the same mathematical description based on fuzzy sets and fuzzy logic (see the book by Dubois and Prade [18] for more information).

The basic notion of the possibility theory is a *possibility distribution function* $\pi(\omega)$, defined on the possibility space Ω (a frame of discernment) which may not be the same as the sample space defined in the theory of probability. The value of $0 \leq \pi(\omega) \leq 1$ represents the measure of possibility of an element ω from the set Ω . It is usually assumed that $\sup\{\pi(\omega) : \omega \in \Omega\} = 1$. A possibility measure of a subset A of Ω is defined as $\pi(A) = \sup\{\pi(\omega) : \omega \in A\}$. There exist many versions (extensions) of the possibility theory, but in all of them the axiom of finite additivity, characteristic for the probability theory, has been replaced by the axiom of *maxitivity*. Let A and B be two events, and $\Pi(A)$ and $\Pi(B)$ be, respectively, their possibilities. Then,

$$\Pi(A \cup B) = \max(\Pi(A), \Pi(B)). \quad (2.1)$$

In his seminal paper [95] Lotfi A. Zadeh proposed to use the formalism of the fuzzy sets theory as the mathematical formalism of the possibility theory. According to this proposal the possibility distribution function that assigns measures of possibility (understood according to the assumed interpretation of this notion) to elements of a certain set (or equivalently, to values of a certain numerical variable) may be interpreted as the membership function assigned to that set. This interpretation allows to use a well developed formal mechanism of the fuzzy sets theory in many different applications. The book by Dubois and Prade [18] describes the links between these two theories, and presents methods for the calculation of numerical values of possibility and necessity measures, both typical for the possibility theory. Moreover, Dubois and Prade [18] pointed out possible links between possibility and probability. According to this interpretation possibility distributions may be regarded as upper envelopes for families of probability distributions. The mutual relation between these two major theories of uncertainty have been later explained and clarified using the Dempster–Shafer theory of evidence. The recent results published in papers of Walley and de Cooman [89], and de Cooman [9] show that measures of possibility are the special case of imprecise probabilities, and thus have a well defined behavioural interpretation.

The original motivation for the development of the possibility theory was to describe imprecise notions or imprecise pieces of information given as statements of a natural language such as, e.g. “costs are high”, “time to failure is about 5 hours”, etc. As a matter of fact, the founders of the possibility theory saw this theory as fundamentally different from the probability theory. They considered the possibility theory as the formalism for the description of uncertain events or uncertain (partial) information in cases where the probability theory failed to provide satisfactory description. On the other hand, the theory of evidence (also known as the theory of belief functions) proposed originally by Dempster [12] and further developed by Shafer [76] aimed at the generalization of the probability theory for dealing with such problems. The basic assumptions of the Dempster–Shafer theory of evidence look similar to the basic assumptions of the theory of probability. It is assumed that there exists a certain possibility space Ω , but probability measures, called in

this theory “probability mass assignments”, in contrast to the probability theory, are defined on its whole power set 2^Ω . This allows to assign probability to an event formed by a set algebra of the elements of the possibility space Ω which in the case of this particular event are indistinguishable, as this is typical for imprecisely described notions.

In the Dempster–Shafer theory of evidence uncertainty is measured using belief functions. A belief function Bel , defined on all subsets of the possibility space Ω , is written in the form

$$Bel(A) = \sum_{B \subseteq A} m(B), \quad (2.2)$$

where m is a probability mass assignment function on all subsets of Ω , such that $m(\emptyset) = 0$, $m(B) \geq 0$ for all $B \subseteq \Omega$ and $\sum_{B \subseteq \Omega} m(B) = 1$. In the Dempster–Shafer theory of evidence there exists also a notion of plausibility which is conjugate to the notion of belief. The conjugate function to the belief function is called the plausibility function Pl , and is defined by

$$Pl(A) = 1 - Bel(A^C) = \sum_{B \cap A \neq \emptyset} m(B), \quad (2.3)$$

where A^C is the complement of the set A . It has been shown that there exists a close relationship between the Dempster–Shafer theory of evidence and the possibility theory. When all elements, say A_1, A_2, \dots, A_n , of the set Ω form a nested set (such that $A_1 \subseteq A_2 \subseteq \dots \subseteq A_n$) then there exists a direct relationship between the mass probability assignments of the Dempster–Shafer theory and the possibility distribution defined as a membership function of a certain fuzzy set (see [19] for more general results). Thus, the possibility theory (for one of its possible interpretations) may be regarded as a special case of the more general Dempster–Shafer theory of evidence. This relationship was used by several authors, who proposed methods for making probability – possibility transformations. Unfortunately, a unique one-to-one transformation between probability and possibility does not exist. Thus, any transformation of this type can lead to loss of information (especially from less precise, but containing more information about the event of interest, possibility to more precise probability), and transformation methods proposed by some authors differ in the methodology used to decrease that loss. One of these methods, proposed by Klir [29, 48] is based on the principle of information invariance. The other approach, based on the optimization of information content, has been proposed by Dubois et al. [21]. More information on the problem of probability – possibility transformation can be also found in [20].

Despite their very close relations, the possibility theory and the Dempster–Shafer theory of evidence are used in different areas of application. The Dempster–Shafer theory is mainly used in building computer expert systems or, in a more general setting, in computerized decision support systems. It is not used, however, in data analysis and in those instances of decision-making processes where statistical

data (precise or imprecise) have to be merged with subjective information. In this particular domain of application the possibility theory is used in practice much more frequently. We will discuss those applications of the possibility theory in the next section of this paper.

2.2.4 *Imprecise Probabilities and Their Generalizations*

Possibility theory and its main tool, the possibility distribution, have been found very useful for the formal description of imprecise information. However, claims – expressed, for example, by Zadeh – that it is also useful for the formalization of imprecise descriptions of probabilities (in statements like “event A is much more probable than event B ”) have been questioned by Walley [86]. In one of his toy examples Walley considers the following information about possible outcomes (win(W), draw (D) or loss(L)) of a football game [86, 87]:

- (a) Probably *not* W .
- (b) W is more probable than D .
- (c) D is more probable than L .

Walley convincingly explains that information of this type cannot be used, without making some arbitrary assumptions, for the precise evaluation of the probability of, e.g. the win $P(W)$. The only consistent evaluation can be done in terms of imprecise probabilities: lower probability $\underline{P}(W)$ and upper probability $\bar{P}(W)$.

Different concepts, such as *interval probabilities* or *non-additive probabilities*, introduced independently by many authors, are covered by the framework of imprecise probabilities. The most comprehensive theory of imprecise probabilities, defined as lower and upper probabilities, was proposed by Walley [86] who introduced the notion of coherent lower (upper) probability. Coherent lower probability may be interpreted as a lower envelope of a set of probability measures that fulfills certain coherence requirements. The similar interpretation exists for the upper probability. The behavioural interpretation of lower probabilities was proposed by Walley [86]. This interpretation is based on the generalization of a similar interpretation of subjective probabilities introduced by de Finetti. According to Walley (see also [87]) the lower (or upper) probability of an event A can be interpreted by specifying acceptable betting rates for betting on (or against) A . If the betting odds on A are x to $1 - x$ one will bet *on* A if $x \leq \underline{P}(A)$ and *against* A if $x \geq \bar{P}(A)$. The choice is not determined if x is between $\underline{P}(A)$ and $\bar{P}(A)$. The basic properties of lower and upper probabilities can be summarized as follows:

- (a) $\underline{P}(\emptyset) = \bar{P}(\emptyset) = 0$.
- (b) $\underline{P}(\Omega) = \bar{P}(\Omega) = 1$.
- (c) $\bar{P}(A) = 1 - \underline{P}(A^C)$.
- (d) $0 \leq \underline{P}(A) \leq \bar{P}(A) \leq 1$.
- (e) $\underline{P}(A) = \underline{P}(B) \leq \underline{P}(A \cup B) \leq \underline{P}(A) + \bar{P}(B) \leq \bar{P}(A \cup B) \leq \bar{P}(A) + \bar{P}(B)$, for disjoint events A and B .
- (f) $\underline{P}(A \cup B) + \underline{P}(A \cap B) \geq \underline{P}(A) + \underline{P}(B)$, for all events A and B .

It is worth to note, that precise probabilities are the special case of lower and upper probabilities such that $\underline{P}(A) = \bar{P}(A)$. Moreover, possibility and necessity measures of the possibility theory and belief and plausibility measures of the Dempster–Shafer theory of evidence are lower and upper probabilities. On the other hand, as it was noted by Walley (see, e.g. [87] and [88]) not all lower and upper probabilities can be interpreted as possibility measures or measures of the Dempster–Shafer theory.

In certain problems of decision-making lower and upper probabilities are not sufficient for dealing with imprecise information. Their further generalization was proposed by Walley (see [86,88]) in his theory of lower and upper previsions. These measures of uncertainty are introduced in terms of *gambles*. A gamble X , defined on a non-empty space Ω of the outcomes of an experiment represents an uncertain reward $X(\omega)$ if the outcome of the experiment is $\omega \in \Omega$. Gambles are expressed in units of some linear utility scale. Below, we present the formal definition of lower and upper previsions, as it was given in [88].

Definition 2.1 (Walley [88]). A bounded mapping from Ω to \mathbb{R} (the real numbers) is called a *gamble*. Let \mathcal{K} be a nonempty set of gambles. A mapping $\underline{P} : \mathcal{K} \rightarrow \mathbb{R}$ is called a *lower prevision* or *lower expectation*. A lower prevision is said to be *coherent* when it is the lowest envelope of some set of linear expectations, i.e. when there is a nonempty set of probability measures, \mathcal{M} , such that $\underline{P}(X) = \min\{E_P(X) : P \in \mathcal{M}\}$ for all $X \in \mathcal{K}$, where $E_P(X)$ denotes the expectation of X with respect to P . The conjugate upper prevision is determined by $\bar{P}(X) = -\underline{P}(-X) = \max\{E_P(X) : P \in \mathcal{M}\}$.

The lower prevision $\underline{P}(X)$ can be interpreted as a supremum acceptable buying price for X . Indicators of events are particular cases of gambles. In such a case, lower and upper previsions coincide with lower and upper probabilities. More information about lower (and upper) previsions can be found in the cited above papers of Walley or in the survey paper by Miranda [61].

The lower (upper) previsions seem to be general enough for the description of subjectively perceived uncertainty. Classical (Bayesian) probabilities, measures of possibility, and Dempster–Shafer measures of evidence can be interpreted as special cases of lower (upper) previsions. From a theoretical point of view this theory is sufficiently well developed. However, there exist some basic problems which require further investigations. For example, the problem of updating the values of imprecise probabilities when new pieces of information are available (i.e. the problem of conditioning) still needs some investigations, as a single generalization of the Bayes updating rule has not been proposed yet. The existing problems with updating procedures are related, for example, to the problems of dealing with observations whose prior probabilities are equal to zero. Other problems arise in relation to concepts of independence or conditional independence. There exist also problems with modeling (in terms of uncertainty) the concepts of preference and weak preference (for example, lower previsions cannot distinguish preference from weak preference). All these problems (for more information, see [88]) motivate researchers to look for more general mathematical models of uncertainty. Some of these models have

been indicated in the first subsection of this section. It is interesting that the need to develop more general models of uncertainty has been also recognized in the community of classical Bayesians. The concept of robust Bayesian inference (see, e.g. the paper by Berger [4]) seems to be closely related to the problems presented in this paper. Moreover, some concepts of frequency-based robust statistics (the notion of ϵ -contamination) can be interpreted using the language of the theory of imprecise probabilities and its generalizations.

2.3 Fuzzy Random Variables and Fuzzy Statistics

The brief description of the existing main theories of uncertainty presented in the preceding section shows that neither of them is fully sufficient to cope with real problems where statistical data are both random and imprecise. The attempts to propose such a theory resulted with the introduction of the notion of a fuzzy random variable. This notion has been defined by many authors. Historically, the first widely accepted definition was proposed by Kwakernaak [54, 55]. Kruse [52] proposed an interpretation of this notion, and according to this interpretation a fuzzy random variable \tilde{Z} may be considered as a perception of an unknown usual random variable $Z : \Omega \rightarrow \mathcal{R}$, called an original of \tilde{Z} . Below, we present another, slightly modified, version of this definition presented in Grzegorzewski [37].

A fuzzy set \tilde{A} described by an upper semicontinuous membership function $\mu_A : \mathcal{R} \rightarrow [0, 1]$ is a fuzzy number when it is:

- (a) Normal, i.e. there exists a $t_0 \in \mathcal{R}$ such that $\mu(t_0) = 1$.
- (b) Fuzzy convex, i.e. $\mu(\lambda s + (1 - \lambda)t) \geq \min\{\mu(s), \mu(t)\}$ for $s, t \in \mathcal{R}$ and $\lambda \in [0, 1]$.
- (c) The set $\{u_0\} = cl\{t \in \mathcal{R} : \mu(t) > 0\}$ is a compact set.

The operator in (c) denoted by cl is the closure operator (see, e.g. Dubois and Prade [15] for a more detailed description). A space of all fuzzy numbers will be denoted by $\mathcal{FN}(\mathcal{R})$. We have $\mathcal{FN}(\mathcal{R}) \subset \mathcal{F}(\mathcal{R})$, where $\mathcal{F}(\mathcal{R})$ denotes a space of all fuzzy sets on the real line. Fuzzy numbers are completely defined by their α -cuts. The α -cut, $\alpha \in (0, 1]$, of a fuzzy number \tilde{A} with its membership function μ_A is a closed crisp set defined as

$$X_\alpha = \{t \in \mathcal{R} : \mu_X(t) \geq \alpha\}. \quad (2.4)$$

In order to describe α -cuts let us use the following notation: $A_\alpha = [A_\alpha^L, A_\alpha^U]$, where

$$A_\alpha^L = \inf\{t \in \mathcal{R} : \mu_A(t) \geq \alpha\}, \quad (2.5)$$

$$A_\alpha^U = \sup\{t \in \mathcal{R} : \mu_A(t) \geq \alpha\}. \quad (2.6)$$

Definition 2.2 (Grzegorzewski [37]). Let (Ω, \mathcal{A}, P) be a probability space, where Ω is a set of all possible outcomes of the random experiment, \mathcal{A} is a σ -algebra of subsets of Ω , and P is a probability measure.

A mapping $\tilde{X} : \Omega \rightarrow \mathcal{FN}(\mathcal{R})$, where $\mathcal{FN}(\mathcal{R})$ is the space of all fuzzy numbers, is called a fuzzy random variable if it satisfies the following properties:

1. $\{X_\alpha(\omega) : \alpha \in [0, 1]\}$ is a set representation of $\tilde{X}(\omega)$ for all $\omega \in \Omega$.
2. For each $\alpha \in (0, 1]$ both $X_\alpha^L = X_\alpha^L(\omega) = \inf X_\alpha(\omega)$ and $X_\alpha^U = X_\alpha^U(\omega) = \sup X_\alpha(\omega)$, are usual real-valued random variables on (Ω, \mathcal{A}, P) .

There exists also another popular definition of a fuzzy random variable proposed by Puri and Ralescu [68] and based on the notion of set-valued mapping and random sets. Below, we present this definition in a form given in [32].

Definition 2.3 (Gil et al. [32]). Let $\mathcal{FN}(\mathcal{R})$ be the space of all fuzzy numbers. Given a probability space (Ω, \mathcal{A}, P) , a mapping $\tilde{X} : \Omega \rightarrow \mathcal{FN}(\mathbb{R}^p)$ is said to be a *fuzzy random variable* (also called *fuzzy random set*) if for all $\alpha \in (0, 1]$ the set-valued mappings $X_\alpha : \Omega \rightarrow \mathcal{K}(\mathbb{R}^p)$, where \mathcal{K} is the class of the non-empty subsets of \mathbb{R}^p , defined so that for all $\omega \in \Omega$ $X_\alpha(\omega) = (X(\omega))_\alpha$, are random sets (that is, Borel-measurable mapping with the Borel σ -field generated by the topology associated with the Hausssdorf metric on $\mathcal{K}(\mathbb{R}^p)$).

Fuzzy random variables may be used to model random and imprecise measurements. First statistical methods for the analysis of such imprecise fuzzy data were developed in the 1980s. Kruse and Meyer [53] proposed a general methodology for dealing with fuzzy random data. In case of their methodology fuzzy random data are described by fuzzy random variables defined according to Definition 2.2. This assumption has very important practical consequences. First of all it means that there exists an underlying non-fuzzy probability distribution that governs the origins of the observed imprecise fuzzy data. The parameters of this distribution have non-fuzzy values, but because of the fuzziness of observed data they cannot be precisely estimated. Their fuzziness comes directly from the fuzziness of statistical data and disappears when statistical data are precise. Therefore, fuzzy statistical methods developed according to the methodology proposed by Kruse and Meyer shall be regarded as straightforward generalization of classical (non-fuzzy) statistical methods. Using the aforementioned methodology Kruse and Meyer [53] proposed methods for the construction of estimators and confidence intervals for the parameters of the probability distributions of fuzzy random variables. According to the interpretation of fuzzy randomness proposed in [53] the observed values of estimators of the parameters of the probability distributions of fuzzy random variables are fuzzy. For this reason these estimators are called fuzzy, despite the fact that their fuzziness is only the reflection of the fuzziness of data, and not of the fuzziness of the estimation procedure itself. The same methodology may be applied to the estimators of the limits of confidence intervals that are also represented by fuzzy numbers. The most important practical consequence of the adoption of the Kruse and Meyer's methodology is that all relevant formulae for fuzzy estimators and other fuzzy statistics can be obtained by fuzzification of well known formulae of classical non-fuzzy statistics. Relevant formulae are obtained by straightforward application of the Zadeh's extension principle (for its definition see the book by Dubois and Prade[15] or any other textbook on fuzzy sets). As this operation is purely

technical, we are not going to present this problem in details. The appropriate formulae can be found in papers devoted to the estimation of parameters of concrete probability distribution when available statistical data are fuzzy. Below, we present only a simple illustrative example.

Let us consider the problem of the estimation of the parameter θ of the exponential distribution defined by the density function $f(t) = (1/\theta)\exp(-t/\theta)$, $t > 0$. The maximum likelihood estimator of this parameter is given as a simple arithmetic average of the observed values in the sample, i.e.

$$\theta^* = \sum_{i=1}^n t_i/n, \quad (2.7)$$

where $t_i, i = 1, \dots, n$ are the values observed in the sample. Suppose now, that instead of crisp values $t_i, i = 1, \dots, n$ we observe fuzzy values described by their respective α -cuts $[t_{i,L}^\alpha, t_{i,U}^\alpha], i = 1, \dots, n, \alpha \in (0, 1]$. Now, we can apply the Zadeh's extension principle and fuzzify (2.7) arriving at the fuzzy version of θ^* described by the set of respective α -cuts:

$$\left[\sum_{i=1}^n t_{i,L}^\alpha/n, \sum_{i=1}^n t_{i,R}^\alpha/n \right] \quad (2.8)$$

The same approach may be applied in all cases when there exist explicit formulae for computation of point estimators and confidence intervals in the case of crisp data. When these formulae do not exist (for example, when maximum likelihood estimators are obtained as numerical solutions of nonlinear equations) the application of the Zadeh's extension principle requires the solution of complex mathematical programming problems.

Despite the fact that the generalization of well known statistical methods to the fuzzy case is relatively straightforward, the construction of fuzzy statistical tests and making statistical decisions is far from being trivial. Fuzzy statistical tests may be developed for testing both non-fuzzy (precise) and fuzzy (imprecise) statistical hypotheses, and for fuzzy (imprecise) and non-fuzzy (precise) statistical data. For example, statistical methods for testing fuzzy hypotheses have been considered in the papers by Saade and Schwarzlander [73], Saade [72], Watanabe and Imaizumi [90], Arnold [1, 2] Taheri and Behboodian [80], and Grzegorzewski and Hryniewicz [39]. When the data are also fuzzy, interesting solutions have been proposed in the papers by Arnold [1], Casals et al. [6], Kruse and Meyer [53], Saade [72], Saade and Schwarzlander [73], Son et al. [79], Watanabe and Imaizumi [90], Römer and Kandel [70], and Montenegro et al. [62]. Grzegorzewski [37] has proposed a unified approach for testing statistical hypotheses with vague data which is a direct generalization of the classical approach. Below, we present his definition of the fuzzy statistical test.

Let $\tilde{Z}_1, \dots, \tilde{Z}_n$ denote a fuzzy sample, i.e. a sample consisting of fuzzy random variables representing fuzzy perception of the usual random sample Z_1, \dots, Z_n

from the population described by the probability distribution P_{Θ} , and let δ be a given number from the interval $(0, 1)$. Grzegorzewski [37] has defined a fuzzy test as follows:

Definition 2.4 (Grzegorzewski [37]). A function

$$\varphi : (\mathcal{FN}(\mathcal{R}))^n \rightarrow \mathcal{F}(\{0, 1\}), \quad (2.9)$$

where $\mathcal{F}(\{0, 1\})$ is the set of possible decisions, is called a fuzzy test for the hypothesis H , on the significance level δ , if

$$\sup_{\alpha \in [0, 1]} P \{ \omega \in \Omega : \varphi_{\alpha}(\tilde{Z}_1(\omega), \dots, \tilde{Z}_n(\omega)) \subseteq \{0\} | H \} \leq \delta$$

where φ_{α} is the α -level set (α -cut) of $\varphi(\tilde{Z}_1, \dots, \tilde{Z}_n)$.

When we test statistical hypotheses about the values of the parameters of probability distributions we utilize a well known equivalence between the set of values of the considered probability distribution parameter for which the null hypothesis is accepted and a certain confidence interval for this parameter. The same equivalence exists in the case of statistical tests with fuzzy data.

When statistical data are precise (crisp), for testing the null hypothesis $H : \theta \leq \theta_0$, and the alternative hypothesis $K : \theta > \theta_0$ we use a one-to-one correspondence between the acceptance region for this test on the significance level δ and the one-sided confidence interval $[\underline{\pi}_l, +\infty)$ for the parameter θ on the confidence level $1 - \delta$, where $\underline{\pi}_l = \underline{\pi}_l(Z_1, \dots, Z_n; \delta)$. This equivalence was utilized by Kruse and Meyer [53] who introduced the notion of a fuzzy confidence interval for the unknown parameter θ , when the data are fuzzy. In the considered case, a fuzzy equivalent of $[\underline{\pi}_l, +\infty)$ can be defined by the following α -cuts (for all $\alpha \in (0, 1]$):

$$\begin{aligned} \underline{\Pi}_{\alpha}^L &= \underline{\Pi}_{\alpha}^L(\tilde{Z}_1, \dots, \tilde{Z}_n; \delta) \\ &= \inf \left\{ t \in \mathcal{R} : \forall i \in \{1, \dots, n\} \exists z_i \in (\tilde{Z}_i)_{\alpha} \right. \\ &\quad \left. \text{such that } \underline{\pi}_l(z_1, \dots, z_n; \delta) \leq t \right\} \end{aligned} \quad (2.10)$$

Similarly, we can define a fuzzy equivalent of the one-sided confidence interval $(-\infty, \bar{\pi}_u]$, as given in Grzegorzewski [37]:

$$\begin{aligned} \bar{\Pi}_{\alpha}^U &= \bar{\Pi}_{\alpha}^U(\tilde{Z}_1, \dots, \tilde{Z}_n; \delta) \\ &= \sup \left\{ t \in \mathcal{R} : \forall i \in \{1, \dots, n\} \exists z_i \in (\tilde{Z}_i)_{\alpha} \right. \\ &\quad \left. \text{such that } \bar{\pi}_u(z_1, \dots, z_n; \delta) \geq t \right\} \end{aligned} \quad (2.11)$$

where $\bar{\pi}_u(z_1, \dots, z_n; \delta) = \underline{\pi}_l(z_1, \dots, z_n; 1 - \delta)$.

The notion of the one-sided fuzzy interval can be used to define a fuzzy test. In the considered case of one-sided statistical hypothesis, a function $\varphi : (\mathcal{FN}(\mathcal{R}))^n \rightarrow \mathcal{F}(\{0, 1\})$ with the following α -cuts:

$$\varphi_\alpha(\tilde{Z}_1, \dots, \tilde{Z}_n) = \begin{cases} \{1\} & \text{if } \theta_0 \in (\underline{\Pi}_\alpha \setminus (\neg \underline{\Pi})_\alpha), \\ \{0\} & \text{if } \theta_0 \in ((\neg \underline{\Pi})_\alpha \setminus \underline{\Pi}_\alpha), \\ \{0, 1\} & \text{if } \theta_0 \in (\underline{\Pi}_\alpha \cap (\neg \underline{\Pi})_\alpha), \\ \emptyset & \text{if } \theta_0 \notin (\underline{\Pi}_\alpha \cup (\neg \underline{\Pi})_\alpha) \end{cases} \quad (2.12)$$

is a fuzzy test for $H : \theta \leq \theta_0$, against $K : \theta > \theta_0$, on the significance level δ (Grzegorzewski [37]). In a similar way, we can define fuzzy tests for testing other one-sided hypotheses such as $H : \theta \geq \theta_0$, against $K : \theta < \theta_0$, and for testing two-sided hypotheses about θ .

It is worthy to note that in certain cases the application of the fuzzy test defined above does not lead to a clearly indicated decision. This feature is far from being unexpected because, unless we make some additional assumptions, we should not expect precise answers to questions presented in a form of statistical hypotheses, if we infer these answers from the analysis of imprecise data. Let us note, however, that we face the similar situation when we use classical statistical methods. In that case a decision cannot be made without setting in advance an appropriate significance level of test. In case of fuzzy statistical data the knowledge of the significance level is not enough, so we have to use some additional indicators that would be helpful in making decisions. There exist several approaches that are suitable for solving this problem. One of these approaches which is formulated in the language of the possibility theory has been proposed by Hryniewicz [43].

In order to present a possibilistic approach to the problem of statistical testing when both data and statistical hypotheses are imprecise let us consider a fuzzy equivalent of testing the hypothesis $H : \vartheta \leq \vartheta_H$ when we observe a random sample (X_1, \dots, X_n) . In case of precisely formulated hypotheses and precise statistical data we can use a well known method (for reference, see, e.g. the book of Lehmann [57]) and calculate a one-sided confidence interval on a confidence level $1 - \delta$ from the formula $[\pi_L(X_1, \dots, X_n; 1 - \delta), \infty)$. We reject the null hypothesis on the significance level δ if the observed value of $\pi_L(X_1, \dots, X_n; 1 - \delta)$ is larger than ϑ_H , i.e. when the inequality $\vartheta_H < \pi_L(x_1, \dots, x_n; 1 - \delta)$ holds. Similarly, we reject the hypothesis $H : \vartheta \geq \vartheta_H$ on the significance level δ when the inequality $\vartheta_H > \pi_U(x_1, \dots, x_n; 1 - \delta)$ holds, where $\pi_U(x_1, \dots, x_n; 1 - \delta)$ is the observed value of the upper limit of the one-sided confidence interval $(-\infty, \pi_U(X_1, \dots, X_n; 1 - \delta)]$ on a confidence level $1 - \delta$. When we test the hypothesis $H : \vartheta = \vartheta_H$ on the significance level δ we reject it if either $\vartheta_H < \pi_L(x_1, \dots, x_n; 1 - \delta/2)$ or $\vartheta_H > \pi_U(x_1, \dots, x_n; 1 - \delta/2)$ holds, where $\pi_L(x_1, \dots, x_n; 1 - \delta/2)$ is the observed in the sample value of the lower limit of the two-sided confidence interval $\pi_L(X_1, \dots, X_n; 1 - \delta/2)$ on a confidence level $1 - \delta$, and the observed value of its upper limit $\pi_U(X_1, \dots, X_n; 1 - \delta/2)$ is given by $\pi_U(x_1, \dots, x_n; 1 - \delta/2)$. Thus,

when we test a hypothesis about the value of the parameter ϑ we find a respective confidence interval, and compare it to the hypothetical value.

Dubois et al. [22] proposed to use statistical confidence intervals of parameters of probability distributions for the construction of possibility distributions of these parameters in a fully objective way. According to their approach, the family of two-sided confidence intervals

$$[\pi_L(x_1, \dots, x_n; 1 - \delta/2), \pi_U(x_1, \dots, x_n; 1 - \delta/2)], \delta \in (0, 1) \quad (2.13)$$

forms the *possibility distribution* $\tilde{\vartheta}$ of the estimated value of the unknown parameter ϑ . In a similar way it is possible to construct one-sided possibility distributions based on one-sided nested confidence intervals. Hryniewicz [43] proposed to compare this possibility distribution with a hypothetical value of the tested parameter. For this purpose he proposed to use the necessity of strict dominance measure introduced by Dubois and Prade [16] for measuring the necessity of the strict dominance relation $\tilde{A} \succ \tilde{B}$, where \tilde{A} and \tilde{B} are fuzzy sets. This measure, called the *Necessity of Strict Dominance* index (*NSD*), is defined as

$$NSD = Ness(\tilde{A} \succ \tilde{B}) = 1 - \sup_{x, y; x \leq y} \min\{\mu_A(x), \mu_B(y)\}. \quad (2.14)$$

Hryniewicz [43] has shown that in the classical case of precise statistical data and precisely defined statistical hypotheses the value of the *NSD* index is equal to the *p*-value of the test.

In case of fuzzy data the confidence intervals used for the construction of the possibility distribution of the estimated parameter θ can be replaced by their fuzzy equivalents, calculated according to the methodology proposed by Kruse and Meyer [53]. In his paper Hryniewicz [43] assumes that the value of the significance level of the corresponding statistical test δ is equal to the possibility degree α that defines the respective α -cut of the possibility distribution of $\tilde{\vartheta}$. He also assumes that in the possibilistic analysis of statistical tests on the significance level δ we should take into account only those possible values of the fuzzy sample whose possibility is not smaller than δ . Thus, the α -cuts of the membership function $\mu_F(\vartheta)$ denoted by $[\mu_{F,L}^{(\alpha)}, \mu_{F,U}^{(\alpha)}]$ are equivalent to the α -cuts of the respective fuzzy confidence intervals on a confidence level $1 - \alpha$.

In order to consider the most general case let us also assume that the hypothetical value of the tested parameter may be also imprecisely defined by a fuzzy number $\tilde{\theta}_H$ described by the membership function $\mu_H(\theta)$. Possibilistic evaluation of the results of statistical fuzzy test consists now in the comparison of the possibility distribution of the estimated parameter θ , and the possibility distribution of the hypothetical value of this parameter. Let us illustrate this procedure by assuming that our fuzzy hypothesis is given as $H : \theta \leq \tilde{\theta}_H$. In the case of crisp data we compare the lower limit of the one-sided confidence interval on a given confidence level $1 - \delta$ with the respective α -cut of the membership function that describes $\tilde{\theta}_H$. In the possibilistic framework described in [43] it means that we compare the possibility distribution

of $\tilde{\vartheta}_L$ based on the left-hand sides of the confidence intervals with the fuzzy value of $\tilde{\theta}_H$. This distribution is defined by the membership function

$$\mu_{F,L}^{(\alpha)} = \inf_{\gamma \geq \alpha} \Pi_L^\gamma(\tilde{X}_1, \dots, \tilde{X}_n; 1 - \delta), \quad (2.15)$$

where

$$\begin{aligned} \Pi_L^\gamma &= \Pi_L^\alpha(\tilde{X}_1, \dots, \tilde{X}_n; 1 - \delta) \\ &= \inf \left\{ t \in \mathcal{R} : \forall i \in \{1, \dots, n\} \exists x_i \in (\tilde{X}_i)_\gamma \right. \\ &\quad \left. \text{such that } \pi_L(x_1, \dots, x_n; 1 - \delta) \leq t \right\}, \end{aligned} \quad (2.16)$$

and $\pi_L(x_1, \dots, x_n; 1 - \delta)$ is the left-hand side limit of the classical confidence interval. In a similar way we can define a fuzzy equivalent of the upper limit of the one-sided confidence interval $(-\infty, \pi_U(X_1, \dots, X_n; 1 - \delta)]$.

In the presence of fuzzy data we have to compare the possibility distribution $\tilde{\vartheta}_{F,L}$ of the estimated value of the unknown parameter θ represented by its α -cuts given by (2.15) with the fuzzy value of $\tilde{\theta}_H$. In such a case we have to find the intersection point of the membership function $\mu_{F,L}(\vartheta)$ and the left-hand side of $\mu_H(\theta)$. The *NSD* index of the relation $\tilde{\vartheta}_{F,L} > \tilde{\theta}_H$ is equal to one minus the ordinate of this point, i.e.

$$Ness(\tilde{\vartheta}_{F,L} > \tilde{\theta}_H) = 1 - \sup \min(\mu_{F,L}(\vartheta), \mu_H(\theta)). \quad (2.17)$$

The *NSD* index defined by (2.17) can be regarded as the generalization of the observed test size p (also known as *p-value* or *significance*) for the case of imprecisely defined statistical hypotheses and vague statistical data. In exactly the same way we can find the *NSD* index for other one-sided and two-sided statistical hypotheses.

Statistical analysis of fuzzy random data can be also done in the Bayesian framework. First results presenting the Bayesian decision analysis for imprecise data were given in papers by Casals et al. [6, 7], and Gil [30]. In these papers the authors described fuzzy observations using the notion of the fuzzy information system by Zadeh [95] and Tanaka et al. [82]. As this approach seems to be not very effective in practical applications, other approaches have been proposed, for example, by Viertl [84], Frühwirth-Schnatter [28], and Taheri and Behboodian [81]. Comprehensive Bayesian model comprising fuzzy data, fuzzy hypotheses, and fuzzy utility function has been proposed in the paper by Hryniewicz [42].

When we interpret fuzzy random variables according to the definition proposed by Puri and Ralescu (see Definition 2.3 above) the statistical analysis of fuzzy data is unfortunately not so simple. The reasons for this difficulty stem from the fact that in that case the underlying classical probability distribution does not exist anymore. For example, it is difficult to formulate fuzzy equivalents of the Central Limit Theorem, as the concept of asymptotic normal distribution cannot be directly applied. Thus, the statistical procedures have to be constructed on a different

theoretical basis. Some authors, see [32] for an overview of the problem, define statistical tests in terms of distances, in different metrics, between the observed fuzzy value of a test statistics and the hypothetical value of a certain characteristic of the fuzzy random variable, e.g. its fuzzy expected value. Therefore, the resulting statistics, such as a variance of the fuzzy random quantity, are real-valued in contrast to fuzzy statistics arriving from the model by Kwakernaak and Kruse. From among few papers devoted to this problem one can mention the papers by Körner [50], Körner and Näther [51], and Montenegro et al. [62]. The problem of lacking underlying probability distribution can be overcome by using a bootstrap methodology, as it has been recently proposed, in the papers by Gil et al. [33], González-Rodríguez et al. [35], and Montenegro et al. [63]. In the case of the Bayesian analysis of fuzzy random variables interesting results have been proposed in the paper by Gil and López-Díaz [31].

2.4 Applications of Fuzzy Statistics in Systems Analysis

Systems analysis is oriented on solving complex problems where precise mathematical models are used for a simplified (and sometimes even oversimplified) description of reality. The main problem of every researcher who has to apply the methods of systems analysis in real applications is related to coping with uncertainties of different kinds. What is important, not all of these uncertainties can be described by well developed methods like theory of probability and mathematical statistics. The methodology of fuzzy statistics, presented in the previous section, gives possibility to describe phenomena where probabilistic randomness is merged with possibilistic imprecision (fuzziness). In this section we present two applications of this methodology which seem to be useful in solving real problems. The character of this paper does not allow us to present too many details that may be necessary to fully understand these applications. The details will be presented in forthcoming papers dedicated to particular problems.

2.4.1 *Example 1: Verification of the Kyoto Protocol*

First, let us consider the problem of the verification of commitments agreed in the Annex I to Kyoto Protocol. The Parties who accept the Kyoto Protocol agreed to reduce the national emissions of greenhouse gases by a specified percentage. The main problem with the verification of these commitments stems from a fact that these emissions cannot be directly measured. Therefore, according to the IPCC Guidance document [46], the total emission X is estimated as a sum of emissions of every type of activity, evaluated indirectly using certain measures describing those activities. For example, the emission from electric coal power plants is evaluated using the knowledge about the amount of burned coal. A simple, but commonly

used in practice, model has a linear form

$$x_i = \sum_{j=1}^m x_{ij} = \sum_{j=1}^m c_{ij} a_{ij}, i = 0, 1, \dots, \quad (2.18)$$

where a_{ij} is the j -th activity measure for the i -th year, and c_{ij} is the emission factor for the i -th year that enables us to calculate greenhouse gas emission knowing the activity measure for the j -th activity. At a national scale the values of activity measures a_{ij} are definitely uncertain, and the level of associated uncertainty strongly depends on the type of activity. Similarly, the values of emission factors c_{ij} may be highly uncertain, as in many cases they are evaluated using experts opinions. More detailed information on this problem can be found in papers by Winiwarter [94], and by Rypdal and Winiwarter [71].

Suppose now, that for each activity we observe a time series, which consists of $n + 1$ yearly observations: $(a_{0j}, a_{1j}, \dots, a_{nj})$, $j = 1, \dots, m$. For sake of simplicity we assume that these are realizations of stochastic processes, and the uncertainty related to their values are purely random. The nature of uncertainty assigned to the associated emission factors $(c_{0j}, c_{1j}, \dots, c_{nj})$, $j = 1, \dots, m$, is hardly easy for precise evaluation. This uncertainty contains undoubtedly a random factor (for example, for an electric coal power plant the emission rate varies randomly with randomly varying quality of burned coal), but may also contain another factor, related to imprecise opinions of experts. For this reason specialists assume that the values of emission factors, and more generally, the values of emissions from different activities, should be evaluated in terms of intervals of possible values. We claim, however, that even in case of very vague information about the values of c_{ij} this assumption is too restrictive. It seems to us that the representation of this information using possibility distributions is much more informative. Therefore, we assume that each value of the emission factors is given as a *fuzzy number* represented by a set of its α -cuts: $[c_{ij,L}^\alpha, c_{ij,R}^\alpha]$, $\alpha \in (0, 1]$. Therefore, the evaluated yearly emission is a realization of a *fuzzy random variable* defined as

$$\tilde{X}_i = \sum_{j=1}^m \tilde{X}_{ij} = \sum_{j=1}^m \tilde{c}_{ij} A_{ij}, i = 0, 1, \dots, \quad (2.19)$$

where A_{ij} is a random value of the j -th activity in the i -th year.

Suppose now, that having the observed realizations of fuzzy random variables \tilde{x}_{ij} , $j = 1, \dots, m$, $j = 0, 1, \dots, n$ we want to predict the total emission for the commitment year k . Let $(a_{kj,L}^\gamma, a_{kj,R}^\gamma)$ be the prediction confidence interval on the confidence level γ for the amount of the j -th activity in the commitment year k . Note, that for $\gamma = 0$ this interval shrinks to the point-wise forecast for the value of x_{kj} . From the theory presented in the previous section we know that *fuzzy confidence interval* for \tilde{X}_{kj} on the confidence level γ is represented by a set of α -cuts (nested α -level intervals): $(a_{kj,L}^\gamma * c_{ij,L}^\alpha, a_{kj,R}^\gamma * c_{ij,R}^\alpha)$. The construction of the fuzzy

prediction interval for the total emission X_k in the commitment year k is a difficult task and, in general, may require the application of Monte Carlo methods. However, when the forecasts for a_{kj} are described by normal distributions this construction is straightforward.

According to the Annex I to Kyoto Protocol a country fulfills its commitment if in the commitment year k its emission does not exceed the value $y_k = \rho x_0$, where x_0 is the emission in the base year, and ρ is a coefficient agreed upon in the Kyoto Protocol. Our analysis of the verification problem we begin from the simplest case when the value of y_k is given, and the activity measures in the commitment year a_{kj} , $j = 1, \dots, m$ have been already established. Thus, we have to verify if $\tilde{x}_k \leq y_k$. From the theory of fuzzy sets we know that the unique method for the verification of this inequality does not exist. However, for this purpose we can use the *NSD* index defined by (2.14) for the relation $y_k \succ \tilde{x}_k$. By simple calculations we can show that this possibility measure is greater than zero if the α -cut of \tilde{x}_k at the level $\alpha = 1$ is located to the left of y_k , i.e. if the inequality $x_{k,R}^1 \leq y_k$ holds. Thus, we have

$$NSD = 1 - \alpha_* \quad (2.20)$$

where

$$\alpha_* : x_{k,R}^{\alpha_*} = y_k. \quad (2.21)$$

From these equations we can also see that $NSD = 1$ if the whole support of \tilde{x}_k is located to the left of y_k . In a more general case, when the emission in the base year is given as a fuzzy number (due to the fuzziness of the emission factors), the commitment may be verified by the calculation of the *NSD* index for the relation $\tilde{y}_k \succ \tilde{x}_k$. In such a case, for the computation of *NSD* we use (2.20), but the value of α_* is now calculated from the following expression:

$$\alpha_* : x_{k,R}^{\alpha_*} = y_{k,L}^{\alpha_*}. \quad (2.22)$$

In the most general case we may want to know in advance, after the evaluations of emissions in years numbered from 0 (base year) to n have been observed, if the commitment in the year k is likely to be fulfilled. We can solve this problem by the verification of a statistical hypothesis that the expected value of the predicted emission (estimated from fuzzy random data) is lower than a given fuzzy valued limit. Thus, we have the problem described in the previous section. The *NSD* index that describes the necessity that this hypothesis is true can be calculated from (2.15)–(2.17).

2.4.2 Example 2: Sequential Testing of a Hypothesis About the Mean Value in the Normal Distribution

As a possible application of the fuzzy statistical methodology in the systems analysis we may consider the problem of testing a hypothesis about the mean value of a random variable described by a normal distribution when sampling costs are high,

and we are forced to observe as few sample items as possible. This situation often happens when we have to control costs of a large project, which consists of many individually assessed partial costs. If the number of partial costs is relatively large, we can assume – following the Central Limit Theorem – that the observed total cost, say X , is distributed according to the normal distribution $N(\mu, \sigma)$. Suppose now that we are interested in keeping the total costs constant for a certain period of time at a level μ_0 . A simple, and the most effective statistical test for verification of the statistical hypothesis $H_0 : \mu = \mu_0$ against the alternative $H_1 : \mu = \mu_1$ is the sequential probability ratio test proposed originally by Wald (see the book by Lehmann [57] for more information) for the case of the known value of the standard deviation σ . The test statistic, based on a sample $(x_1 \dots x_n)$ is, in that case, a simple sum of re-scaled observations

$$S_n = \sum_{i=1}^n (x_i / \sigma). \quad (2.23)$$

Let α be the probability of the type-I error (probability of an erroneous rejection of H_0), and β be the probability of the type-II error (probability of an erroneous acceptance of H_1). We accept the null hypothesis H_0 if

$$S_n \geq \frac{A}{\mu_1 - \mu_0} + \frac{n}{2}(\mu_1 + \mu_0), \quad (2.24)$$

where $A = \beta/(1 - \alpha)$. We reject H_0 in favour of H_1 if

$$S_n \leq \frac{B}{\mu_1 - \mu_0} + \frac{n}{2}(\mu_1 + \mu_0), \quad (2.25)$$

where $B = (1 - \beta)/\alpha$. If neither of these inequality holds, we have to increase the sample size by one, and repeat the same procedure.

In the sequential test described above we have assumed that the value of σ is known. In practice, we never know this value in advance, but when the amount of historical data is large enough we can estimate σ , and take this estimated value as the known one. However, when the available amount of data is scarce, as it is usually the case in the analysis of large systems, we cannot proceed this way. A possible way out is to use a procedure proposed by Hryniewicz [44] for the analysis of reliability data.

Having some historical data we can estimate the value of σ . When the available sample size is equal to m , and the estimated value is given by σ_m^* , we can calculate the two-sided confidence interval for the unknown value of σ :

$$\left(\sigma_L(\gamma) = \sqrt{\frac{(m-1)(\sigma_m^*)^2}{\chi_{m-1}^2((1+\gamma)/2)}}, \sigma_R(\gamma) = \sqrt{\frac{(m-1)(\sigma_m^*)^2}{\chi_{m-1}^2((1-\gamma)/2)}} \right), \quad (2.26)$$

where $\chi_{m-1}^2(\gamma)$ is the quantile of order γ of the chi-square distribution with $m - 1$ degrees of freedom. The confidence intervals defined by (2.26) can be used for the construction of the *possibility distribution* of σ defined by its δ -cuts (we use here the symbol of δ because the symbol α , traditionally used in this context, has been used for the description of the type-I error of the sequential test). The left-hand side limits of this possibility distribution are given by

$$\mu_L^\delta(\sigma) = \begin{cases} \sigma_L(1 - \delta) & \text{if } \delta \geq \delta_0 \\ \sigma_L(1 - \delta_0) & \text{if } \delta < \delta_0 \end{cases}, \quad (2.27)$$

where δ_0 is a small number close to 0 (e.g. 0.01). Similarly, the right-hand limits are given by

$$\mu_R^\delta(\sigma) = \begin{cases} \sigma_R(1 - \delta) & \text{if } \delta \geq \delta_0 \\ \sigma_R(1 - \delta_0) & \text{if } \delta < \delta_0 \end{cases}. \quad (2.28)$$

If we assume that σ is a fuzzy number defined by this possibility distribution we immediately find that the test statistic S_n becomes also fuzzy. Thus, we cannot directly verify if inequalities (2.24) and (2.25) are fulfilled. However, we may assume that they are fulfilled if the *NSD* index for respective fuzzy relations is greater than a prescribed value.

In this second example of the application of fuzzy statistics we haven't used any subjective imprecise fuzzy information. The fuzziness has been introduced in a purely objective way using some historical statistical data. We can also generalize this problem by allowing imprecise hypotheses about the values of μ_0 and μ_1 . The methodology for dealing with such a problem is the same, but the necessary calculations become much more complicated.

2.5 Conclusions

In the paper we have presented different methods used for mathematical modeling of uncertainty. We have concentrated on modeling of epistemic uncertainty, i.e. uncertainty evaluated subjectively by human beings like experts which reflects lack of full information about phenomena of interest. We have presented basic information about the most important (among many others) theories: theory of probability, which is the basic theory of uncertainty, and alternative theories of uncertainty like Dempster–Shafer theory of evidence, theory of imprecise probabilities, and theory of possibility. Considerable space has been devoted to the theory of fuzzy random variables which links classical probability with fuzzy sets and theory of possibility. This choice stems from the fact that, as for now, this is the only theory which provides mathematical foundations for comprehensive analysis of data representing aleatoric randomness and epistemic imprecision.

It has to be stressed once more that the theory of probability (together with classical mathematical statistics) seems to be the only reasonable methodology for dealing with randomness. However, its ability to model other types of uncertainty

has been questioned by many scholars. Alternative theories of uncertainty, like those mentioned above, have been developed to cope with problems where classical probability provides questionable solutions. Some of these theories, like theory of imprecise probabilities, may be regarded as generalizations of the classical probability, some others, like theory of possibility, were introduced as completely different theories. It has to be noted, however, that recently published theoretical results reveal many links between all these theories making them rather complementary than competitive.

Alternative theories of uncertainty are relatively very young in comparison to the theory of probability. Therefore, they are not as matured as the theory of probability, and there exist questions for which unequivocal answers have not been proposed yet. The current situation is well illustrated by an experiment proposed in the paper by Oberkampf et al. [66]. In this experiment prominent representatives of different methodologies were asked to provide numerical evaluation of outputs of systems described by equations with uncertain parameters. Different types of sources of uncertainty were represented in this experiment, allowing for using different types of models, and different methods for aggregation of uncertainty. The results of this experiment, described in [23], show that the usage of different approaches resulted in different evaluations. It is interesting, however, that prominent representatives of the probabilistic approach, who were invited to this experiment, either refused to provide their evaluation, arguing that the input information was unrealistically imprecise, or provided results obtained after the introduction of additional assumptions which had not been stated in the original description of this experiment.

The results of the experiment mentioned above, and analyses presented in overview papers, like the paper by Helton et al. [40] or by Möller and Beer [65], reveal the most important challenges specialists in dealing with uncertainty are confronted with. First, in case of the analysis of complex systems and structures the analysts who apply alternative approaches for coping with uncertainty face very serious computational problems. They have to use methods, such as interval computations, which are computationally much more demanding in comparison to methods used in case of traditional probabilistic models. Second, Monte Carlo methodology for simulation of systems has to be extended in order to cope with information which is not random by nature. For example, the existing methodology for simulation of fuzzy random variables is still under development. Third, for some alternative theories of uncertainty unequivocally accepted methods of propagation of uncertainty have not been proposed yet. And finally, some very general theories of uncertainty still need methods for coping with statistical data.

The challenges mentioned above are less visible when we apply classical theory of probability. The price for this favorable situation is sometimes high. We have to introduce additional assumptions, which in presence of partial information (or partial ignorance) cannot be verified. Therefore, there is still a need for finding more appropriate methods for coping with uncertainty. The methodology of fuzzy random variables, described in numerous recently published papers, and books like the books by Viertl [85] or by Möller and Beer [64], in our opinion seems to be, as for now, the most useful in practice.

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Chapter 3

On the Approximation of a Discrete Multivariate Probability Distribution Using the New Concept of t -Cherry Junction Tree

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Abstract Most everyday reasoning and decision making is based on uncertain premises. The premises or attributes, which we must take into consideration, are random variables, so that we often have to deal with a high dimensional discrete multivariate random vector. We are going to construct an approximation of a high dimensional probability distribution that is based on the dependence structure between the random variables and on a special clustering of the graph describing this structure. Our method uses just one-, two- and three-dimensional marginal probability distributions. We give a formula that expresses how well the constructed approximation fits to the real probability distribution. We then prove that every time there exists a probability distribution constructed this way, that fits to reality at least as well as the approximation constructed from the Chow–Liu dependence tree. In the last part we give some examples that show how efficient is our approximation in application areas like pattern recognition and feature selection.

3.1 Introduction

The goal of our paper is to approximate a high dimensional joint probability distribution using two and three-dimensional marginal distributions, only.

The idea of such kind of approximations was given by Chow and Liu [7]. In their work they construct a first order tree taking into account the mutual information gains of all pairs of random variables. They proved that their approximation is optimal in the sense of Kullback–Leibler divergence [12, 15].

In order to give an approximation that uses lower dimensional marginal probability distributions, there are many algorithms developed. Most of them first construct

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a Bayesian network (directed acyclic graph) see [1] and then they obtain from this in several steps a junction tree. See [11] for a good overview.

Many algorithms were developed for obtaining a Bayesian Network. These algorithms start with the construction of the Chow–Liu tree and then this graph is transformed, by adding edges and then delete the superfluous edges using conditional independence tests. The number of conditional independence tests is diminished by searching the minimal d -separating set [2, 6].

After the graphical structure is determined a number of quantitative operations have to be performed on it (see [13] and [9]).

In our paper we suppose to be known just the three-dimensional marginals of the joint probability distribution (indeed that implies that the second and first order marginals are known, too). From this information we first construct a graphical model, and then a junction tree, named t -cherry-junction tree.

The construction of our graphical model is inspired by the graphical structure, named cherry tree, and t -cherry tree introduced by Bukszár and Prékopa in [3].

We emphasize that our method does not use the construction of the Bayesian network first, we are going to use just the third order marginals and the information contents related to them, and to certain pairs of random variables involved.

The paper is organized as follows:

- The second section contains the theory, formulas, and connections between them that are used in the third section.
- The third section contains the introduction of the concept of the t -cherry-junction tree, the formula that gives the Kullback–Leibler divergence associated to the approximation introduced, and a proof that the approximation associated to the t -cherry-junction tree is at least as good as Chow–Liu’s approximation is.
- The last section contains some applications of the approximation introduced in order to exhibit some advantages of this approach.

3.2 Preliminaries

3.2.1 Notations

Let $\mathbf{X} = (X_1, X_2, \dots, X_n)^T$ be an n -dimensional random vector with the joint probability distribution

$$P(X_1 = x_{i_1}^1, \dots, X_n = x_{i_n}^n), i_1 = 1, \dots, m_1, \dots, i_n = 1, \dots, m_n.$$

For this we will use also the abbreviation

$$P(\mathbf{X}) = P(X_1, X_2, \dots, X_n).$$

This shorter form will be applied in sums and products, too. So we will write

$$\sum_{\mathbf{X}} P(\mathbf{X}) = \sum_{i_1=1}^{m_1} \cdots \sum_{i_n=1}^{m_n} P(X_1 = x_{i_1}^1, \dots, X_n = x_{i_n}^n) (= 1) \quad (3.1)$$

and for example if $H = \{j, k, l\}$ is a three element subset of the index (vertex) set $\{1, 2, \dots, n\}$ then $\mathbf{X}_H = (X_j, X_k, X_l)^T$ and

$$\sum_{\mathbf{X}_H} P(\mathbf{X}_H) = \sum_{i_j=1}^{m_j} \sum_{i_k=1}^{m_k} \sum_{i_l=1}^{m_l} P(X_j = x_{i_j}^j, X_k = x_{i_k}^k, X_l = x_{i_l}^l) (= 1) \quad (3.2)$$

and

$$\prod_{\mathbf{X}_H} P(\mathbf{X}_H) = \prod_{i_j=1}^{m_j} \prod_{i_k=1}^{m_k} \prod_{i_l=1}^{m_l} P(X_j = x_{i_j}^j, X_k = x_{i_k}^k, X_l = x_{i_l}^l) \quad (3.3)$$

$P_a(\mathbf{X})$ will denote the approximating joint probability distribution of $P(\mathbf{X})$.

3.2.2 Cherry Tree and t -Cherry Tree

The cherry tree is a graph structure introduced by Bukszár and Prékopa (see [3]). A generalization of this concept, called hyper cherry tree can be found in [4] by Bukszár and Szántai. Let be given a nonempty set of vertices V .

Definition 3.1. The graph defined recursively by the following steps is called *cherry tree*:

1. Two vertices connected by an edge is the smallest cherry tree.
2. By connecting a new vertex of V , by two new edges to two existing vertices of a cherry tree, one obtains a new cherry tree.
3. Each cherry tree can be obtained from (1) by the successive application of (2).

Remark 3.1. A cherry tree is an undirected graph. We emphasize that it is not a tree.

Definition 3.2. We call *cherry*, a triplet of vertices formed from two existing vertices and a new one connected with them in step (2) of Definition 3.1.

For the cherry we will use the notation introduced in [4] by Bukszár and Szántai: $(\{l, m\}, k)$, where l and m are the existing vertices, k is the newly connected vertex and $\{k, l\}, \{k, m\}$ are the new edges.

Remark 3.2. From a set of n vertices we obtain a cherry tree with $n - 2$ cherries.

Remark 3.3. We denote by \mathcal{H} the set of all cherries of the cherry tree and by ϵ the set of edges of the cherry tree.

Remark 3.4. A cherry tree is characterized by a set V of vertices, the set \mathcal{H} of cherries and the set ϵ of edges. We denote a cherry-tree by $\Delta = (V, \mathcal{H}, \epsilon)$.

In our paper we need the concept of the t -cherry tree introduced by Bukszár and Prékopa in [3]. To get the concept of t -cherry tree one has to apply the more restrictive Step (2') in Definition 3.1 instead of Step (2):

(2') By connecting a new vertex of V by two new edges to two *connected* vertices of a cherry tree, one obtains a new cherry tree.

Definition 3.3. The graph defined recursively by (1), (2') and (3) is called *t -cherry tree*.

Remark 3.5. A pair of adjacent vertices from the t -cherry tree may be used several times, for connecting new vertices to them.

3.2.3 Junction Tree

The junction tree is a very prominent and widely used structure for inference in graphical models (see [5] and [12]).

Let $X = \{X_1, \dots, X_n\}$ be a set of random variables defined over the same probability field and $\mathbf{X} = (X_1, \dots, X_n)^T$ an n -dimensional random vector.

Definition 3.4. A tree with the following properties is called junction tree over X :

1. To each node of the tree, a subset of X called cluster and the marginal probability distribution of these variables is associated.
2. To each edge connecting two clusters of the tree, the subset of X given by the intersection of the connected clusters and the marginal probability distribution of these variables is associated.
3. If two clusters contain a random variable, than all clusters on the path between these two clusters contain this random variable (running intersection property).
4. The union of all clusters is X .

Notations:

- \mathcal{C} – the set of clusters
- C – a cluster
- \mathbf{X}_C – the random vector with the random variables of C as components
- $P(\mathbf{X}_C)$ – the joint probability distribution of \mathbf{X}_C
- \mathcal{S} – the set of separators
- S – a separator
- \mathbf{X}_S – the random vector with the random variables of S as components
- $P(\mathbf{X}_S)$ – the joint probability distribution of \mathbf{X}_S

The junction tree provides a joint probability distribution of \mathbf{X} :

$$P(\mathbf{X}) = \frac{\prod_{C \in \mathcal{C}} P(\mathbf{X}_C)}{\prod_{S \in \mathcal{S}} P(\mathbf{X}_S)^{(v_S - 1)},}$$

where v_S is the number of those clusters which contain all of the variables involved in S .

3.3 t -Cherry-Junction Tree

3.3.1 Construction of a t -Cherry-Junction Tree

Let $X = \{X_1, \dots, X_n\}$ be a set of random variables defined over the same probability field and denote by $\mathbf{X} = (X_1, \dots, X_n)^T$ the corresponding n -dimensional random vector. Together with the construction of the t -cherry tree over the set of vertices $V = \{1, \dots, n\}$ we can construct a t -cherry-junction tree in the following way.

Algorithm 1. *Construction of a t -cherry-junction tree.*

1. The first cherry of the t -cherry tree identifies the first cluster of the t -cherry junction tree. The vertices of the first cherry give the indices of the random variables belonging to the first cluster.
2. Similar way each new cherry of the t -cherry tree identifies a new cluster of the t -cherry-junction tree. The separators contain pairs of random variables with indices corresponding to the connected vertices used in Step (2') of Definition 3.3.

Theorem 3.1. *The t -cherry-junction tree constructed by Algorithm 1 is a junction tree.*

Proof. We can check the statements of Definition 3.4:

- The first statement of definition is obvious.
- The separator that connects two clusters contains the intersection of the two clusters. This follows from the construction of the separator sets.
- Let us suppose that there exists a variable X_m which belongs to two clusters on a path, so that on this path exist a cluster that does not contain X_m . This implies that on this path exist two neighboring cluster so that one of them contains X_m and the other one does not contain X_m . This means that in the t -cherry tree there are two cherries connected so that one of them contains the vertex m the other one does not contain the vertex m . According to the point (2) of Definition 3.1, m is a new vertex connected, but since X_m belongs already to another cluster, the vertex m belongs to another cherry, too. That is a contradiction, because that means that m is not a new vertex.

- The union of all sets associated to the clusters is X , because of the union of set of indices is V .

Remark 3.6. To each cluster $\{X_l, X_m, X_k\}$ a three-dimensional joint probability distribution $P(X_l, X_m, X_k)$ can be associated. To each separator $\{X_l, X_m\}$ a two-dimensional joint probability distribution $P(X_l, X_m)$ can be associated.

The joint probability distribution associated to the t -cherry-junction tree is a joint probability distribution over the variables X_1, \dots, X_n , given by

$$P(\mathbf{X}) = \frac{\prod_{\{X_l, X_m, X_k\} \in \mathcal{C}} P(X_l, X_m, X_k)}{\prod_{\{X_l, X_m\} \in \mathcal{S}} P(X_l, X_m)^{v_{lm}-1}}, \quad (3.4)$$

where $v_{lm} = \#\{\{X_l, X_m\} \subset C \mid C \in \mathcal{C}\}$.

3.3.2 *The Approximation of the Joint Distribution Over X by the Distribution Associated to a t -Cherry-Junction Tree*

Chow and Liu introduced a method to approximate optimally an n -dimensional, discrete joint probability distribution by the one- and two-dimensional probability distributions using first-order dependence tree. It is shown that the procedure presented in their paper yields an approximation with minimum difference of information, in the sense of Kullback–Leibler divergence.

In this part, we first give a formula for the Kullback–Leibler divergence between the approximated distribution associated to the t -cherry-junction tree and the real joint probability distribution; we then conclude what we have to take into account to minimize the divergence. For the proof of this theorem we need a lemma:

Lemma 3.1. *In a t -cherry-junction tree for each variable $X_m \in X$:*

$$\#\{\{X_l, X_m\} \mid (\{X_l, X_m\}, X_k) \in \mathcal{C}\} = \#\{(\{X_l, X_m\}, X_k) \mid (\{X_l, X_m\}, X_k) \in \mathcal{C}\} - 1$$

Proof. Let denote $t = \#\{\{X_l, X_m\} \mid (\{X_l, X_m\}, X_k) \in \mathcal{C}\}$ for a given $X_m \in X$.

- Case $t = 0$.

The statement is a consequence on one hand of Definition 3.4 that is the union of all sets associated to the nodes (clusters) is X , so every vertex from X , have to appear at least in one cluster. On the other hand, if X_m would be contained in two clusters than there must exist one separator set containing this vertex (point 3) in Definition 3.4), but we supposed $t = 0$.

- Case $t > 0$.

If two clusters contain a variable X_m , than all clusters from the path between the two clusters contain X_m (running intersection property). From this results that

the clusters containing X_m are the nodes of a connected graph, and this graph is a first order tree. If this graph contains $t + 1$ clusters, than it contains t separator sets (definition of a tree in which we have separators instead of edges and clusters instead of nodes). From the definition of the junction tree (Definition 3.4) results that every separator set connecting two clusters contains the variables from the intersection of the clusters. So there are t separator sets that contain the variable X_m .

The goodness of the approximation of a probability distribution can be quantified by the Kullback–Leibler divergence between the real and the approximating probability distributions (see for example [8] or [15]). The Kullback–Leibler divergence expresses somehow the distance between two probability distributions. As smaller value it has as better the approximation is.

Theorem 3.2. *If we denote by $P_a(\mathbf{X})$ the approximating joint probability distribution associated to the t -cherry-junction tree [see formula (3.4)], then the Kullback–Leibler divergence between the real and the approximating joint probability distribution is given by:*

$$\begin{aligned}
 I(P(\mathbf{X}), P_a(\mathbf{X})) &= -H(\mathbf{X}) - \sum_{(X_l, X_m, X_k) \in \mathcal{C}} I(X_l, X_m, X_k) \\
 &+ \sum_{\{X_l, X_m\} \in \mathcal{S}} (v_{lm} - 1) I(X_l, X_m) + \sum_{k=1}^n H(X_k).
 \end{aligned} \tag{3.5}$$

Proof.

$$\begin{aligned}
 I(P(\mathbf{X}), P_a(\mathbf{X})) &= \sum_{\mathbf{X}} P(\mathbf{X}) \log_2 \frac{P(\mathbf{X})}{P_a(\mathbf{X})} \\
 &= \sum_{\mathbf{X}} P(\mathbf{X}) \log_2 P(\mathbf{X}) - \sum_{\mathbf{X}} P(\mathbf{X}) \log_2 P_a(\mathbf{X}) \\
 &= -H(\mathbf{X}) - \sum_{\mathbf{X}} P(\mathbf{X}) \log_2 \frac{\prod_{\{X_l, X_m, X_k\} \in \mathcal{C}} P(X_l, X_m, X_k)}{\prod_{\{X_l, X_m\} \in \mathcal{S}} P(X_l, X_m)^{v_{lm}-1}} \\
 &= -H(\mathbf{X}) - \sum_{\mathbf{X}} P(\mathbf{X}) \left[\log_2 \prod_{\{X_l, X_m, X_k\} \in \mathcal{C}} P(X_l, X_m, X_k) \right. \\
 &\quad \left. - \log_2 \prod_{\{X_l, X_m\} \in \mathcal{S}} P(X_l, X_m)^{v_{lm}-1} \right] \\
 &= -H(\mathbf{X}) - \sum_{\mathbf{X}} P(\mathbf{X}) \log_2 \prod_{\{X_l, X_m, X_k\} \in \mathcal{C}} P(X_l, X_m, X_k) \\
 &\quad + \sum_{\mathbf{X}} P(\mathbf{X}) \log_2 \prod_{\{X_l, X_m\} \in \mathcal{S}} P(X_l, X_m)^{v_{lm}-1}
 \end{aligned}$$

From Definition 3.4 follows that the union of the clusters of the junction tree is the set X . From the Lemma 3.1 we know that each vertex appears once more in clusters than in separator sets. So by adding and subtracting the sum

$$\sum_{\mathbf{X}} P(\mathbf{X}) \log_2 \left[\prod_{\{X_l, X_m, X_k\} \in \mathcal{C}} P(X_l) P(X_m) P(X_k) \right]$$

we obtain the following:

$$\begin{aligned} & I(P(\mathbf{X}), P_a(\mathbf{X})) \\ &= -H(\mathbf{X}) - \sum_{\mathbf{X}} P(\mathbf{X}) \log_2 \frac{\prod_{\{X_l, X_m, X_k\} \in \mathcal{C}} P(X_l, X_m, X_k)}{\prod_{\{X_l, X_m, X_k\} \in \mathcal{C}} P(X_l) P(X_m) P(X_k)} \\ &+ \sum_{\mathbf{X}} P(\mathbf{X}) \log_2 \frac{\prod_{\{X_l, X_m\} \in \mathcal{S}} P(X_l, X_m)^{v_{lm}-1}}{\prod_{\{X_l, X_m\} \in \mathcal{S}} [P(X_l) P(X_m)]^{v_{lm}-1}} - \sum_{\mathbf{X}} P(\mathbf{X}) \log_2 \prod_{X_k \in X} P(X_k) \\ &= -H(\mathbf{X}) - \sum_{\mathbf{X}} P(\mathbf{X}) \sum_{\{X_l, X_m, X_k\} \in \mathcal{C}} \log_2 \frac{P(X_l, X_m, X_k)}{P(X_l) P(X_m) P(X_k)} \\ &+ \sum_{\mathbf{X}} P(\mathbf{X}) \sum_{\{X_l, X_m\} \in \mathcal{S}} (v_{lm} - 1) \log_2 \frac{P(X_l, X_m)}{P(X_l) P(X_m)} + \sum_{X_k \in X} H(X_k). \end{aligned}$$

Since (X_l, X_m, X_k) and (X_l, X_m) are components of the random vector \mathbf{X} , we have the relations:

$$\begin{aligned} & \sum_{\mathbf{X}} P(\mathbf{X}) \sum_{\{X_l, X_m, X_k\} \in \mathcal{C}} \log_2 \frac{P(X_l, X_m, X_k)}{P(X_l) P(X_m) P(X_k)} \\ &= \sum_{\{X_l, X_m, X_k\} \in \mathcal{C}} \sum_{(X_l, X_m, X_k)^T} P(X_l, X_m, X_k) \log_2 \frac{P(X_l, X_m, X_k)}{P(X_l) P(X_m) P(X_k)} \end{aligned}$$

and

$$\begin{aligned} & \sum_{\mathbf{X}} P(\mathbf{X}) \sum_{\{X_l, X_m\} \in \mathcal{S}} (v_{lm} - 1) \log_2 \frac{P(X_l, X_m)}{P(X_l) P(X_m)} \\ &= \sum_{\{X_l, X_m\} \in \mathcal{S}} \sum_{(X_l, X_m)^T} (v_{lm} - 1) P(X_l, X_m) \log_2 \frac{P(X_l, X_m)}{P(X_l) P(X_m)}. \end{aligned}$$

Taking into account these relations and applying the notion of the mutual information content for two and three variables:

$$I(X_l, X_m) = \sum_{(X_l, X_m)^T} P(X_l, X_m) \log_2 \frac{P(X_l, X_m)}{P(X_l)P(X_m)},$$

$$I(X_l, X_m, X_k) = \sum_{(X_l, X_m, X_k)^T} P(X_l, X_m, X_k) \log_2 \frac{P(X_l, X_m, X_k)}{P(X_l)P(X_m)P(X_k)}$$

we obtain (3.5) and the statement of the theorem has been proved.

Observation 1. We can observe that for minimizing the Kullback–Leibler divergence between the real probability distribution and the approximation obtained from the t -cherry-junction tree we have to maximize the difference between the sum of the mutual divergence of the clusters and the sum of the mutual divergence of the separators denoted by S :

$$S = \sum_{\{X_l, X_m, X_k\} \in \mathcal{C}} I(X_l, X_m, X_k) - \sum_{\{X_l, X_m\} \in \mathcal{S}} (v_{lm} - 1)I(X_l, X_m)$$

Observation 2. If we wish to compare two approximations of a joint probability distribution associated to two different t -cherry-junction trees, we have just to:

- Sum the information contents of the clusters
- Sum the information contents of the separators
- Make the difference between them
- Claim a t -junction-cherry tree be better than another one, if it produces greater value of S

3.3.3 *The Relation Between the Approximations Associated to the First-Order Dependence Tree and t -Cherry-Junction Tree*

A natural question is: can the t -cherry-junction tree give better approximation than the first order tree given by Chow and Liu does? Let us remind that Chow and Liu introduced a method for finding an optimal first order tree that minimizes the Kullback–Leibler divergence.

If $pa(j)$ denotes the parent node of j , the joint probability distribution associated to the Chow–Liu first order dependence tree is given as follows:

$$P_{Ch-L}(\mathbf{X}) = \prod_{i=1}^n P(X_{m_i} | X_{pa(m_i)}),$$

where $\{m_1, \dots, m_n\}$ is a permutation of the numbers $1, 2, \dots, n$ and if $pa(j)$ is the empty set, then by definition $P(X_j | X_{pa(j)}) = P(X_j)$.

In the following Algorithm 2 we will show how one can construct a t -cherry-junction tree from a Chow–Liu first order dependence tree. After this in Theorem 3.3 we prove that the t -cherry-junction tree constructed this way gives at least as good approximation of the real probability distribution as the Chow–Liu first order dependence tree does.

Algorithm 2. *The construction of a t -cherry-junction tree from a Chow–Liu first order dependence tree.*

Let us regard the spanning tree behind the Chow–Liu first order dependence tree. It is sufficient to give an algorithm for constructing a t -cherry tree from this spanning tree. Then by Algorithm 1 we can assign a t -cherry-junction tree to this t -cherry tree:

1. The first cherry of the t -cherry tree let be defined by any three vertices of the spanning tree which are connected by two edges.
2. We add a new cherry to the t -cherry tree by taking a new vertex of the spanning tree adjacent to the so far constructed t -cherry tree.
3. We repeat step (2) till all vertices from the spanning tree become included in the t -cherry tree.

Theorem 3.3. *If $P_{Ch-L}(\mathbf{X}) = \prod_{i=1}^n P(X_{m_i} | X_{pa(m_i)})$ is the approximation associated to the Chow–Liu first order dependence tree there always exists a t -cherry-junction tree with associated probability distribution $P_{t-ch}(\mathbf{X})$ that approximates $P(\mathbf{X})$ at least as well as $P_{Ch-L}(\mathbf{X})$ does.*

Proof. We construct the t -cherry-junction tree from the Chow–Liu first order dependence tree using Algorithm 2.

The Kullback–Leibler divergence formally looks like it is given in (3.5).

In the case of the approximation obtained by the Chow–Liu method, the Kullback–Leibler divergence is given as follows (see [7]):

$$I(P(\mathbf{X}), P_{Ch-L}(\mathbf{X})) = -H(\mathbf{X}) - \sum_{i=1}^n I(X_{m_i}, X_{pa(m_i)}) + \sum_{i=1}^n H(X_i). \quad (3.6)$$

Because the first and last terms of the Kullback–Leibler divergences are the same in the case of the two approximations (3.5) and (3.6), we denote the sums that we have to compare by S_{Ch-L} in the case of Chow–Liu’s approximation and S_{t-ch} in the case of t -cherry-junction tree approximation:

$$S_{Ch-L} = \sum_{X_{m_i} \in X} I(X_{m_i}, X_{pa(m_i)}) \quad (3.7)$$

and

$$S_{t-ch} = \sum_{\{X_l, X_m, X_k\} \in \mathcal{C}} I(X_l, X_m, X_k) - \sum_{\{X_l, X_m\} \in \mathcal{S}} (v_{lm} - 1) I(X_l, X_m) \quad (3.8)$$

In the case of formula (3.7) we can apply the formula

$$I(X, Y) = H(X) - H(X|Y)$$

(see [8], Formula (2.43) on p. 16), while in the case of formula (3.8) we can apply

$$I(X, Y, Z) = H(Z) + I(X, Y) - H(Z|X, Y)$$

which easily can be derived from formulae given in book [8].

So from (3.7) we get

$$\begin{aligned} S_{Ch-L} &= \sum_{X_{m_i} \in X} [H(X_{m_i}) - H(X_{m_i}|X_{pa(m_i)})] \\ &= \sum_{X_{m_i} \in X} H(X_{m_i}) - \sum_{X_{m_i} \in X} H(X_{m_i}|X_{pa(m_i)}) \end{aligned} \quad (3.9)$$

and from (3.8) we get

$$\begin{aligned} S_{t-ch} &= \sum_{\{X_l, X_m, X_k\} \in \mathcal{C}} [H(X_k) + I(X_l, X_m) - H(X_k|X_l, X_m)] \\ &\quad - \sum_{\{X_l, X_m\} \in \mathcal{S}} (\nu_{lm} - 1)I(X_l, X_m) \\ &= \sum_{X_k \in X} H(X_k) - \sum_{\{X_l, X_m, X_k\} \in \mathcal{C}} H(X_k|X_l, X_m) \end{aligned} \quad (3.10)$$

From (3.9) and (3.10) we conclude that we have to compare only

$$\sum_{X_{m_i} \in X} H(X_{m_i}|X_{pa(m_i)})$$

and

$$\sum_{\{X_l, X_m, X_k\} \in \mathcal{C}} H(X_k|X_l, X_m)$$

To each X_k corresponds an X_{m_i} because these both are the vertices of X and each of them appears exactly once. The edge connecting X_{m_i} and $X_{pa(m_i)}$ is contained in the t -cherry tree as it was constructed according to Step (2) in the proof of Theorem 3.3. From this we can conclude that for each X_k contained in a cluster $\{X_l, X_m, X_k\}$ one of X_l and X_m is $X_{pa(k)}$. Now we can apply the inequality (see book [8], Formula (2.130) on p. 36):

$$H(X_k|X_i, X_j) \leq H(X_k|X_i)$$

with equality just in case X_k is conditionally independent of X_j . From this it is obvious that $S_{t-ch} \geq S_{Ch-L}$, and indeed the relation between the Kullback–Leibler

divergences of the two approximations is:

$$I(P(\mathbf{X}), P_{t-ch}(\mathbf{X})) \leq I(P(\mathbf{X}), P_{Ch-L}(\mathbf{X})).$$

This proves the statement of the theorem.

Observation 1. If the underlying dependence structure is a first order dependence tree than we got equality between the two divergences. (Because of the conditional independences that take place between the unlinked vertices of the first order dependence tree).

3.4 Some Practical Results of Our Approximation and Discussions

This section consists of three parts. In the first part we consider two different approximations of a given eight-dimensional discrete joint probability distribution:

- The approximation given by the Chow–Liu method
- The approximation corresponding to the t -cherry tree constructed by the algorithm given in the proof of Theorem 3.3

In the second part of this section we use the approximations to make some prediction, when the values taken by two out of eight random variables are known. From this information we are going to recognize the most probable values of the remaining six random variables. In the third part we use the influence diagram underlying the t -cherry-junction tree to make a feature selection.

Two Approximations of an Eight-Dimensional Discrete Probability Distribution

First we construct an eight-dimensional discrete probability distribution in the following way. The one-dimensional marginal distributions are generated randomly. Then the so called North-West corner algorithm was applied to determine an initial feasible solution of an eight-dimensional transportation problem, where the quantities to be transported were the marginal probability values. This way the “transported probabilities” are concentrated on 44 different directions, i.e. the constructed eight-dimensional discrete probability distribution is concentrated on 44 eight-dimensional vector instead of the possible 2,116,800. An other advantage is that this way we can get as high as possible positive correlations between the components of the random vector \mathbf{X} . For having also high negative correlations, we combined the North-West corners with South-West corners.

Now we suppose that only the two and three-dimensional marginals of the eight-dimensional joint probability distribution are known. Using these we are going to construct a probability distribution associated to the t -cherry-junction tree. First we construct the Chow–Liu spanning tree and then transform the correspondent Chow–Liu dependence tree in a t -cherry junction tree using the Algorithm 2.

Table 3.1 The mutual information gains in decreasing order for the construction of the Chow–Liu spanning tree

Index pair	Information gain
58	1.660755
18	1.644658
28	1.641684
25	1.611500
15	1.605311
56	1.593626
17	1.558543
47	1.556713
16	1.436039
14	1.423012
68	1.421927
26	1.416225
78	1.367303
13	1.334538

Table 3.2 The three variable mutual information contents ordered in decreasing way. The boldfaced ones are used in the t -cherry-junction tree

Index triplet	Information content
158	3.615752
156	3.431799
258	3.414463
168	3.384689
568	3.357803
128	3.322943
125	3.321265
256	3.278311
147	3.276440
178	3.270665
157	3.197210
578	3.167661
268	3.151382
148	3.137220
135	3.120396

The first step is to calculate the mutual information contents of every pair and triplet of random variables. We then order them in descending way.

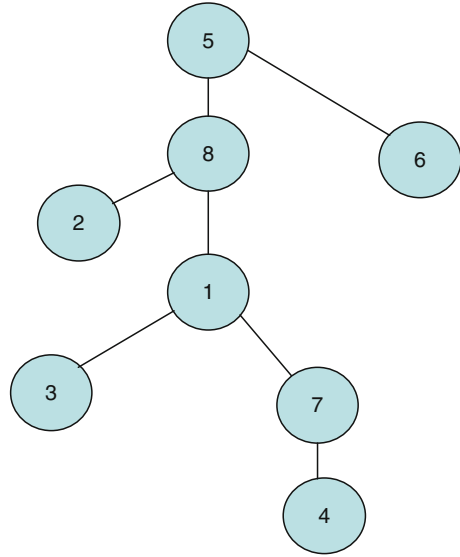
The Chow–Liu tree is constructed by a greedy algorithm from the two-dimensional information gains. In Table 3.1 the ordered mutual information gains are given. The mutual information gains used by Chow–Liu’s method are in boldface. In Fig. 3.1 the Chow–Liu tree can be seen.

The joint probability distribution associated to the constructed Chow–Liu dependence tree is:

Table 3.3 The two variable mutual information contents ordered in decreasing way. The boldfaced ones are used in the t -cherry-junction tree

Index pair	Information content
58	1.660755
18	1.644658
28	1.641684
25	1.611500
15	1.605311
56	1.593626
17	1.558543

Fig. 3.1 The Chow–Liu spanning tree



$$\begin{aligned}
 P_{Ch-L}(\mathbf{X}) &= P(X_5)P(X_8|X_5)P(X_2|X_8)P(X_6|X_5)P(X_1|X_8)P(X_3|X_1)P(X_7|X_1)P(X_4|X_7) \\
 &= \frac{P(X_5X_8)P(X_2X_8)P(X_5X_6)P(X_1X_8)P(X_1X_3)P(X_1X_7)P(X_4X_7)}{P(X_8)P(X_5)P(X_8)P(X_1)P(X_1)P(X_7)}.
 \end{aligned}$$

The Kullback–Leibler divergence corresponding to the divergence between the real probability distribution and the approximation given by Chow–Liu method can be calculated as follows:

$$\begin{aligned}
 I(P(\mathbf{X}), P_{Ch-L}(\mathbf{X})) &= \sum_{i=1}^8 H(X_i) - H(\mathbf{X}) - [I(X_5, X_8) + I(X_2, X_8) + I(X_5, X_6) + I(X_1, X_8) \\
 &\quad + I(X_1, X_3) + I(X_1, X_7) + I(X_4, X_7)] \\
 &= 16.908113 - 4.634363 - 10.990524 = 1.283226.
 \end{aligned}$$

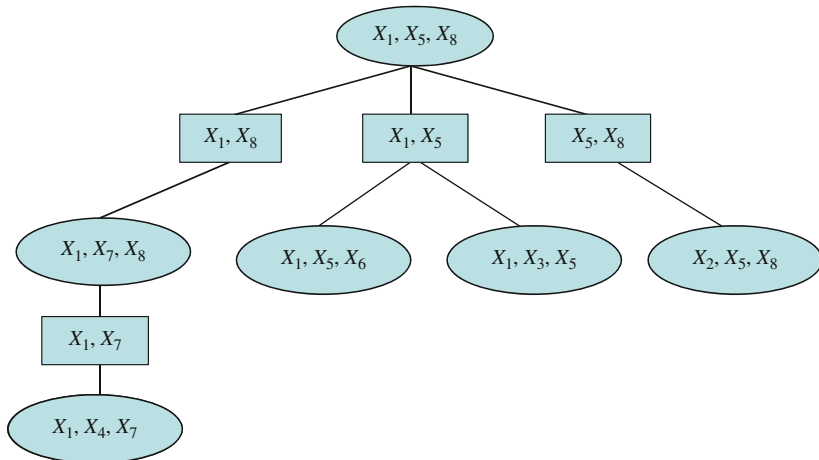


Fig. 3.2 The t -cherry-junction tree

The t -cherry-junction tree constructed by Algorithm 2 can be seen in Fig. 3.2. The three and the two variable mutual information contents corresponding to the clusters and separators of the t -cherry-junction tree are given in bold face in Tables 3.2 and 3.3

The joint probability distribution associated to the constructed t -cherry-junction tree is:

$$P_{t-ch}(\mathbf{X}) = \frac{P(X_1 X_5 X_8) P(X_1 X_7 X_8) P(X_1 X_5 X_6) P(X_1 X_3 X_5) P(X_2 X_5 X_8) P(X_1 X_4 X_7)}{P(X_1 X_8) P(X_1 X_5) P(X_1 X_5) P(X_5 X_8) P(X_1 X_7)}.$$

The Kullback–Leibler divergence between the real probability distribution and the approximation associated to the t -cherry-junction tree is:

$$\begin{aligned} I(P(\mathbf{X}), P_{t-ch}(\mathbf{X})) &= \sum_{i=1}^8 H(X_i) - H(\mathbf{X}) - [I(X_1, X_5, X_8) + I(X_1, X_7, X_8) + I(X_1, X_5, X_6) \\ &\quad + I(X_1, X_3, X_5) + I(X_2, X_5, X_8) + I(X_1, X_4, X_7) - I(X_1, X_8) \\ &\quad - I(X_1, X_5) - I(X_1, X_5) - I(X_5, X_8) - I(X_1, X_7)] \\ &= 16.908113 - 4.634363 - 20.129510 + 8.074578 = 0.218818. \end{aligned}$$

The real eight-dimensional distribution has 44 different vectors with probabilities different from 0. The Chow–Liu approximation has 453 vectors; the t -cherry-junction tree approximation has 93 vectors with probabilities different from zero. From the two Kullback–Leibler divergences calculated above we can observe easily that the approximation associated to the t -cherry-junction tree is much better (“closer” to the reality) than the approximation constructed from the Chow–Liu tree.

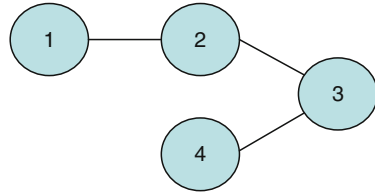
Table 3.4 Comparison of the predicted values with the real ones in the case of the two different approximations

The most probable vectors predicted by Ch-L approx.							$P(\mathbf{X})$	$P_a(\mathbf{X})$	The most probable vectors in the reality							$P(\mathbf{X})$		
7	5	1	7	1	2	7	1	0.003487	0.003443	7	5	1	7	1	2	7	1	0.003487
6	5	1	7	1	2	7	3	0.006802	0.005149	6	5	1	7	1	2	7	3	0.006802
3	3	3	4	4	5	3	5	0.000000	0.001696	3	3	3	5	4	5	5	5	0.009853
2	2	3	4	5	5	3	7	0.011576	0.003718	2	2	3	4	5	5	3	7	0.011576
4	4	3	6	2	3	6	4	0.000000	0.004076	4	4	3	6	3	4	6	4	0.012450
3	2	3	4	4	5	3	6	0.019070	0.008043	3	2	3	4	4	5	3	6	0.019070
8	5	1	7	1	2	7	1	0.019399	0.019155	8	5	1	7	1	2	7	1	0.019399
3	2	3	4	5	5	3	7	0.010207	0.006777	3	2	3	4	4	5	3	7	0.025312
7	5	1	7	1	2	7	2	0.027852	0.027503	7	5	1	7	1	2	7	2	0.027852
9	5	1	7	1	2	8	1	0.034906	0.037489	9	5	1	7	1	2	8	1	0.034906
4	3	3	6	4	5	6	5	0.000000	0.033668	4	3	3	6	4	4	6	5	0.044913
5	4	3	7	2	3	7	4	0.000000	0.030836	5	4	3	7	3	4	7	4	0.049737
1	2	4	1	5	5	2	7	0.027323	0.033325	1	2	4	2	5	5	3	7	0.054170
6	5	3	7	1	2	7	2	0.000000	0.001522	6	5	1	7	1	2	7	2	0.128273
6	4	2	7	3	3	7	4	0.050302	0.009993	6	4	1	7	2	3	7	4	0.152472
The most probable vectors predicted by t-ch approx.							$P(\mathbf{X})$	$P_a(\mathbf{X})$	The most probable vectors in the reality							$P(\mathbf{X})$		
7	5	1	7	1	2	7	1	0.003487	0.003487	7	5	1	7	1	2	7	1	0.003487
6	5	1	7	1	2	7	3	0.006802	0.006802	6	5	1	7	1	2	7	3	0.006802
3	3	3	5	4	5	5	5	0.009853	0.009497	3	3	3	5	4	5	5	5	0.009853
2	2	3	4	5	5	3	7	0.011576	0.009733	2	2	3	4	5	5	3	7	0.011576
4	4	3	6	3	4	6	4	0.012450	0.009917	4	4	3	6	3	4	6	4	0.012450
3	2	3	5	4	5	4	6	0.012682	0.018480	3	2	3	4	4	5	3	6	0.019070
8	5	1	7	1	2	7	1	0.019399	0.019399	8	5	1	7	1	2	7	1	0.019399
3	2	3	4	4	5	3	7	0.025312	0.020359	3	2	3	4	4	5	3	7	0.025312
7	5	1	7	1	2	7	2	0.027852	0.027852	7	5	1	7	1	2	7	2	0.027852
9	5	1	7	1	2	8	1	0.034906	0.034906	9	5	1	7	1	2	8	1	0.034906
4	3	3	6	4	4	6	5	0.044913	0.037797	4	3	3	6	4	4	6	5	0.044913
5	4	3	7	3	4	7	4	0.049737	0.058277	5	4	3	7	3	4	7	4	0.049737
1	2	4	2	5	5	3	7	0.054170	0.044752	1	2	4	2	5	5	3	7	0.054170
6	5	1	7	1	2	7	2	0.128273	0.128273	6	5	1	7	1	2	7	2	0.128273
6	4	1	7	2	3	7	4	0.152472	0.154778	6	4	1	7	2	3	7	4	0.152472

Application for Pattern Recognition

In this part we are testing our approximations for the following pattern recognition problem. We suppose that the values of X_1 and X_8 are known. For these given values we want to predict the most probable values of the other six random variables. In Table 3.4 one can see these predictions made with the help of the approximation.

Fig. 3.3 A possible dependence diagram of four random variables



In the left side of the table the predicted values of the random variables X_2, \dots, X_7 are given for the two different approximations. In the right side of the table the most probable values of the same random variables are given according to the real probability distribution (the same in the case of both approximations). The rows of the table are in ascending order according to the real probabilities. As the wrong predicted values are typed in boldface one easily can find them for each approximation. Let us observe that as long as the number of the wrong predicted values is 13 in the case of the Chow–Liu approximation, the same number equals only 2 in the case of the new t -cherry-junction tree approximation.

Feature Selection: Forecasting the Values of a Random Variable Which Depends on Many Others

The main idea of feature selection is to choose a subset of input random variables by eliminating features with little or no predictive information. In supervised learning the feature selection is useful when the main goal is to find feature subset that produces higher classification accuracy.

In practice many times we have a lot of attributes (random variables) that depend more or less on each other. The problem is how to select a few of them to make a good forecast of the variable we are interested in. The pairwise mutual information contents are not sufficient to make such a decision. To highlight this let us consider the following example. If we have four random variables with the relations between their pairwise mutual information contents: $I(X_2, X_3) > I(X_1, X_3) > I(X_3, X_4)$, and want to take into account only two random variables to forecast the values of X_3 , we would use X_2 and X_1 . But if we have the dependence diagram (see Fig. 3.3) we should decide in an other way. As X_1 influences X_3 only through X_2 we should rather use X_2 and X_4 for the forecast of the values of X_3 . To solve such problems we give a method that uses the t -cherry-junction tree. If we are interested in forecasting a random variable X_i , we have to select from the cherry tree the clusters that contain X_i . We obtain in this way a sub junction tree. This results immediately from the properties of the junction tree. We have to take into consideration, just the variables that belong to this sub junction tree.

If in our earlier numerical example we are interested in the forecast of X_8 we select from the t -cherry-junction tree the clusters that contain X_8 . From Fig. 3.2 these are (X_1, X_5, X_8) , (X_1, X_7, X_8) and (X_2, X_5, X_8) . Now we can conclude that

for our purpose it is important to know the joint probability distribution of the random vector $(X_1, X_2, X_5, X_7, X_8)^T$. This can be obtained as a marginal distribution of the distribution associated to the cherry tree, which can be also expressed as:

$$P(X_1, X_2, X_5, X_7, X_8) = \frac{P(X_1, X_5, X_8)P(X_1, X_7, X_8)P(X_2, X_5, X_8)}{P(X_1, X_8)P(X_5, X_8)}.$$

Now to test our method we calculate the value of the conditional entropy $H(X_8 | X_1, X_2, X_5, X_7) = 0.214946$ from the *real* probability distribution. If we choose another random vector $(X_1, X_2, X_5, X_7, X_8)^T$, then we get $H(X_8 | X_1, X_2, X_5, X_6) = 0.235572$.

It is interesting to see that $I(X_1, X_2, X_5, X_6, X_8) = 7.210594$ is greater than $I(X_1, X_2, X_5, X_7, X_8) = 7.087809$.

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Part II
Robust Solutions Under Uncertainty

Chapter 4

Induced Discounting and Risk Management

T. Ermolieva, Y. Ermoliev, G. Fischer, and M. Makowski

Abstract The goal of this paper is to specify and summarize new approaches to discounting proposed in our catastrophic risk management studies. The main issue is concerned with justification of investments, which may turn into benefits over long and uncertain time horizon. For example, how can we justify mitigation efforts for expected 300-year flood that can occur also next year. The discounting is supposed to impose time preferences to resolve this issue, but this view may be dramatically misleading. We show that any discounted infinite horizon sum of values can be equivalently replaced by undiscounted sum of the same values with random finite time horizon. The expected duration of this stopping time horizon for standard discount rates obtained from capital markets does not exceed a few decades and therefore such rates may significantly underestimate the net benefits of long-term decisions. The alternative undiscounted random stopping time criterion allows to induce social discounting focusing on arrival times of the main concern (stopping time) events rather than horizons of market interests.

In general, induced discount rates are conditional on the degree of social commitment to mitigate risk. Random stopping time events affect these rates, which alter the optimal mitigation efforts that, in turn, change events. This endogeneity of the induced discounting restricts exact evaluations necessary for using traditional deterministic methods and it calls for stochastic optimisation methods. The paper provides insights in the nature of discounting that are critically important for developing robust long-term risk management strategies.

4.1 Introduction

The implication of uncertainties and risks for justifying long-term investments is a controversial issue. How can we justify investments, which may possibly turn into benefits over long and uncertain time horizons in the future? This is a key issue for catastrophic risk management. For example, how can we justify investments in

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climate change mitigations, say, in flood defense systems to cope with foreseen extreme 1,000-, 500-, 250-, and 100-floods? The lack of proper evaluations for dealing with extreme events dramatically contributes to increasing losses from human-made and natural disasters [20]. The analysis of floods that occurred in the summer of 2002 across central Europe [14] shows that the potential areas of vulnerability to extreme floods have multiplied as a consequence of failed development planning. Underestimation and ignorance of low probability/high consequence events have led to the growth of buildings and industrial land and sizable value accumulation in flood prone areas without proper attention being paid to flood mitigations. A challenge is that an endogenously created catastrophe,¹ say a 300-year flood, has never occurred before in a given region. Therefore, purely adaptive policies relying on historical observations provide no awareness of the “unknown” risk although, a 300-year flood may occur next year. For example, the 2002 floods in Austria, Germany and the Czech Republic were classified (in different regions) as 1,000-, 500-, 250-, and 100-year events [14].

A key issue is development of policies with proper long-term perspectives. The traditional discounting is supposed to impose necessary time preferences, but this view may be dramatically misleading. There are several possibilities for choosing discount rates (see, for example, the discussion in [2, 19, 24, 29]). The traditional approach is to use the rates obtained in capital markets. The geometric or exponential discount factor $d_t = (1 + r)^{-t} = e^{-\ln(1+r)t} \approx e^{-rt}$ (for small r) is usually connected with a constant rate r of returns from capital markets. Since returns in capital markets are linked to assets with a lifespan of a few decades, this choice may completely reduce the impacts that investments have beyond these intervals (Sect. 4.2). Another serious problem [21, 31] arises from the use of the expected value $r = E\xi$ and the discount factor $(1 + r)^{-t}$. It implies additional significant reduction of future values in contrast to the expected discount factor $E(1 + \xi)^{-t}$, because $E(1 + \xi)^{-t} \gg (1 + r)^{-t}$. These issues are discussed in Sects. 4.2 and 4.3.

An appropriate interest rate is especially difficult to define when decisions involve time horizons beyond the interests of the current generation. If future generations are not present in the market, e.g., long-term environmental damages are not included in production costs, the market interest rates do not reflect the preferences of future generations. According to Arrow et al. [2] “the observed market rates of interest refer to how individuals are willing to trade off consumption over their own life. These may or may not bear close correspondence to how a society is willing to trade off consumption across generations”.

Debates on proper discount rates for long-term problems have a long-standing history [2, 29]. Ramsey [26] argued that applying a positive discount rate r to discount values across generations is unethical. Koopmans [17], contrary to Ramsey, argued that zero discount rate r would imply an unacceptably low level of current consumption. The use of so-called social discount rates produces two effects [2].

¹ As a consequence of inappropriate policies.

The “prescriptive” approach tends to generate relatively low discount rates and thus favors mitigation measures and the wellbeing of future generations. The “descriptive” approach tends to generate higher discount rates and thus favors less spending on mitigation and the wellbeing of the current generation.

The constant discount rate has only limited justification [3, 12, 24, 29]. As a compromise between “prescriptive” and “descriptive” approaches, Cline [4] argues for a declining discount rate: 5% for the first 30 years, and 1.5% later. There have been proposals for other schedules and attempts to justify the shape of proper decline. Papers [21, 31] show that uncertainty about r produces a certainty-equivalent discount rate, which will generally be declining with time. Weitzman [31] proposed to model discount rates by a number of exogenous time dependent scenarios. He argued for rates of 3–4% for the first 25 years, 2% for the next 50 years, 1% for the period 75–300 years and 0 beyond 300 years. Newel and Pizer [21] analyzed the uncertainty of historical interest rates by using data on the US market rate for long-term government bonds. They proposed a different declining discount rate justified by a random walk model. Chichilinsky [3] proposed a new concept for long-term discounting with a declining discount rate by attaching some weight on the present and the future consumption. All these papers aim to derive an appropriate exogenous social discount rate.

Sections 4.2 and 4.3 develop a different approach for social discounting. It is shown that any discounted sum, so-called net present value (NPV) criterion, $\sum_{t=0}^{\infty} d_t V_t$ of expected values $V_t = E v_t$ for random variables (r.v.) $v_t, t = 0, 1, \dots, d_t = (1 + r_t)^{-t}$ under constant and declining discount rates r_t equals the average undiscounted (in the agreement with Ramsey’s concerns) random sum $E \sum_{t=0}^{\tau} v_t$ with a random stopping time τ defined by the given discounting d_t . Therefore, discount rates can be associated with the occurrences of “stopping time” random events determining a finite “internal” discount-related horizon $[0, \tau]$. The expected duration of τ and its standard deviation σ under modest market interest rates of 3.5% is approximately 30 years, which may have no correspondence with expected, say, 300-year extreme events and $\sigma \approx 300$. Conversely, it is shown that any stopping time random event induces a discounting. A set of mutually exclusive stopping time random events, e.g., 1,000-, 500-, 250-, and 100-year floods, induces discounting with time-declining discount rates. This case corresponds also to the discounting with uncertain discount rates r . In particular, a single stopping time random event with the standard geometric probability distribution induces the standard discounting with constant discount rate r and $d_t = (1 + r)^{-t}$.

The effects of catastrophes on the stream of values $v_t, t = 0, 1, \dots$, differ from the effects of market uncertainties. Section 4.4 indicates that catastrophic events pose new challenges. They often create so-called endogenous, unknown (with the lack and even absence of adequate observations) and interdependent risks, which may potentially affect large territories and communities and, on the other hand, are dramatically affected by risk management decisions. As a consequence, catastrophic risks generally make it impossible to use traditional economic and insurance models [1, 3, 5, 8, 9, 16]. The concept of undiscounted random stopping time criteria

allows to induce social discounting that focuses on arrivals of catastrophic events rather than the lifetime of market products. Since risk management decisions affect the occurrence of disasters in time and space, the induced discounting may depend on spatio-temporal distributions of extreme events and feasible sets of decisions, i.e., it can be viewed as a spatio-temporal discounting. The implicit dependence of the stopping time discounting on random events and decisions calls for the use of stochastic optimization methods, which allows also to address the variability (Remark 4.2) of discounted criteria by using random value $\sum_{t=0}^{\tau} v_t$ even for deterministic $v_t, t = 0, 1, \dots$. Section 4.5 establishes connections of stopping time discounting with dynamic versions of CVaR (Conditional Value-at-Risk) risk measures. Section 4.6 illustrates how misperception of induced discounting may provoke catastrophes. Section 4.7 provides concluding remarks.

4.2 Standard and Stopping Time Induced Discounting

This section illustrated the main idea by using the standard geometric discounting. The choice of discount rate as a prevailing interest rate within a time horizon of existing financial markets is well established [18]. Uncertainties, especially related to extreme events, challenge the possibility of markets to offer proper rates for longer time horizons. The following simple Proposition 4.1 and Remark 4.2 clarify the main concerns.

The traditional financial approaches [18] often use the so-called net present value (NPV) criteria to justify investments. An investment is defined as an expected cash flow stream $V_0, V_1, \dots, V_T, V_t = E v_t$, over a time horizon $T \leq \infty$. Assume that r is a constant prevailing market interest rate, then alternative investments are compared by $V = V_0 + d_1 V_1 + \dots + d_T V_T$, where $d_t = d^t, d = (1 + r)^{-1}, t = 0, 1, \dots, T$, is the discount factor and V denotes NPV.

It is usually assumed that a long-term investment activity has an infinitely long time horizon, i.e.,

$$V = \sum_{t=0}^{\infty} d_t V_t. \quad (4.1)$$

The stream of values $V_t, t = 0, 1, \dots$, can represent an expected cash flow stream of a long-term investment activity. In economic growth models and integrated assessment models [19, 22, 29] the value V_t represents utility $U(x^t)$ of an infinitely living representative agent, or welfare $V_t = \sum_{i=1}^n a_i u_i(x_i^t)$ of a society with representative agents $i = 1, \dots, n$, utilities u_i , consumptions x_i^t and welfare weights a_i . Natural selection theory treats (4.1) as Darwinian fitness [28], where discount factors d_t are associated with hazard rates of an environment (Example 4.2).

The infinite time horizon in (4.1) creates an illusion of truly long-term analysis. Proposition 4.1 shows that in fact deterministic evaluation (4.1) accounts only for values V_t from a finite random horizon $[0, \tau]$ defined by a random stopping time τ with the discount-related geometric probability distribution $P[\tau \geq t] = d_t$.

Proposition 4.1. Consider a discounted sum (4.1) with $d_t = d^t$, $d = (1 + r)^{-1}$, $r > 0$. Let $q = d$, $p = 1 - q$, and τ be a random variable with the geometric probability distribution $P[\tau = t] = pq^t$, $t = 0, 1, \dots$. Then $d_t = P[\tau \geq t]$ and

$$\sum_{t=0}^{\infty} d^t V_t = \sum_{t=0}^{\infty} P[\tau \geq t] V_t = E \sum_{t=0}^{\tau} V_t. \quad (4.2)$$

Conversely, for any stopping time τ with a geometric probability distribution

$$E \sum_{t=0}^{\tau} V_t = \sum_{t=0}^{\infty} d_t V_t, \quad d_t = P[\tau \geq t].$$

Proof. We have $P[\tau \geq t] = \sum_{k=t}^{\infty} pq^k = pq^t(1 - q)^{-1} = q^t = d_t$. Conversely,

$$\begin{aligned} E \sum_{t=0}^{\tau} V_t &= \sum_{t=0}^{\infty} P[\tau = t] \sum_{k=0}^t V_k = \sum_{t=0}^{\infty} pq^t \sum_{k=0}^t V_k \\ &= \sum_{t=0}^{\infty} \left(\sum_{k=t}^{\infty} pq^k \right) V_t = \sum_{t=0}^{\infty} d_t V_t. \end{aligned}$$

That is, any discounted deterministic sum (4.1) equals to the average undiscounted random sum $\sum_{t=0}^{\tau} V_t$ of the same values V_t . In other words, the discount factor $d_t = d^t$ induces an “internal” discount-relate time horizon $[0, \tau]$ with the geometrically distributed τ . Conversely, any geometrically distributed τ and the criterion $E \sum_{t=0}^{\tau} V_t$ induces the geometric discounting in the sum $\sum_{t=0}^{\infty} d_t V_t$.

Remark 4.1. (Random stopping time horizon). We can consider $[0, \tau]$ being a random stopping time horizon associated with the first occurrence of a “killing”, i.e., a catastrophic stopping time event. The probability that this event occurs at $t = 0, 1, \dots$ is p and pq^t is the probability that this event occurs first time at t , i.e., τ has a geometric probability distribution. Since $p = 1 - d$, $d = (1 + r)^{-1}$, then the expected duration of τ , $E\tau = 1/p = 1 + 1/r$. Therefore, for the interest rate of 3.5%, $r = 0.035$, the expected duration is $E\tau \approx 30$ years, i.e., this rate orients the policy analysis on an expected 30-year time horizon. The standard deviation $\sigma = \sqrt{q/p}$, i.e., it equals approximately 30 years. The bias in favor of the present in discounting with the rate of 3.5% is easily illustrated [24]. For a project with long-run benefits or costs, 1 Euro of benefits or costs in years 50, 100, and 200, has a present value respectively of 0.18, 0.003, and practically 0 Euros. Definitely, this rate may have no correspondence to how society has to deal with a 300-year flood, i.e., a flood with the expected arrival time equal to 300 years. Therefore, in the risk management τ can be associated with the arrival of potential catastrophic events rather than with horizons of market interests. The induced social discounting $d_t = P[\tau \geq t]$ in this case would have proper long-term perspectives dependent on spatio-temporal patterns of catastrophes and risk management decisions (see Proposition 4.3 and Sect. 4.4). The discount rate r can be viewed also as a killing (hazard) rate [15] which makes the life expectancy of an otherwise infinitely living

representative agent or society equal to $1 + 1/r$ years. Yet, depending on a concrete situation, stopping time τ can be also associated with the arrival time of a reward.

Remark 4.2. (Variability of NPV). Disadvantages of this standard criterion (4.1) are well known [18]. In particular, the NPV critically depends on the prevailing interest rate which may not be easily defined in practice. In addition, the NPV does not reveal the temporal variability of cash flow streams. Two alternative streams may easily have the same NPV despite the fact that in one of them all the cash is clustered within a few periods, but in another it is spread out evenly over time. This type of temporal heterogeneity is critically important for dealing with catastrophic losses which occur suddenly as a “spike” in time and space [9].

The criterion $E \sum_{t=0}^{\tau} V_t, V_t = E v_t$ has visible advantages. In particular, it allows to address distributional aspects and robust strategies [6] by analyzing the random variable $\sum_{t=0}^{\tau} V_t$ (even for deterministic $v_t = V_t$), e.g., its quantiles defined as maximal $y = y_{\delta}$ satisfying safety constraint

$$P \left[\sum_{t=0}^{\tau} v_t \geq y \right] \geq \delta.$$

Equivalently, y_{δ} maximizes the concave function (see discussion in [6], p. 16)

$$y + \delta^{-1} E \min \left\{ 0, \sum_{t=0}^{\tau} v_t - y \right\}.$$

The optimal value of this function defines the so-called CVaR (Conditional Value-at-Risk) risk measure [27].

Therefore, if variables v_t depend on some decisions x (as in Sect. 4.4), then the maximization of function

$$F(x) = \left[y + \delta^{-1} E \min \left\{ 0, \sum_{t=0}^{\tau} v_t - y \right\} \right].$$

allows easy control of highly nonlinear (even for linear in x function v_t) the safety constraints (quantiles of $\sum_{t=0}^{\tau} v_t$) in an optimal manner defined by a function $F(x)$ that is adjusted to CVaR risk measure (see also Sect. 4.5).

Remark 4.3. (Shock testing). The sensitivity of models w.r.t. “shocks” (extreme scenarios, events, stresses) is often assessed by introducing them into discounted criteria [22, 29]. From Proposition 4.1 it follows that this may lead to serious miscalculations. Let us consider criterion (4.1) with discount factors, $d_t = d^t, d = (1+r)^{-1}$ and assume that a “shock” arrives at a random time moment $\theta \in \{0, 1, \dots\}$ with probability $P[\theta = t] = \pi \gamma^t, \gamma = 1 - \pi = (1 + \rho)^{-1}$. Then the expected value, $E \sum_{t=0}^{\theta} d_t V_t = \sum_{t=0}^{\infty} d^t \gamma^t V_t = E \sum_{t=0}^{\tau} \gamma^t V_t = E \sum_{t=0}^{\min(\tau, \theta)} V_t$, where $P[\tau = t] = pq^t$ with $q = d, p = 1 - q$. Therefore, the stopping time of the “shocked” evaluation $E \sum_{t=0}^{\theta} d^t V_t$ is defined by $\min(\tau, \theta)$. The discount rate of this evaluation is $(1 + r)^{-1} \cdot (1 + p)^{-1} = (1 + r + \rho + r\rho)^{-1}$, i.e., the shocked

evaluation increases the rate of the original discounting and, hence, the bias in favor of the present.

Example 4.1. (Catastrophic Risk Management). The implications of Proposition 4.1 for long-term policy analysis are rather straightforward. Let us consider some important cases. It is realistic to assume [24] that the cash flow stream, typical for investment in a new nuclear plant, has the following average time horizons. Without a disaster the first six years of the stream reflect the costs of construction and commissioning followed by 40-years of operating life when the plant is producing positive cash flows and, finally, a 70-year period of expenditure on decommissioning. The flat discount rate of 5%, as Remark 4.1 shows, orients the analysis on a 20-year time horizon. It is clear that a lower discount rate places more weight on distant costs and benefits. For example, the explicit treatment of a potential 200-year disaster would require at least the discount rate of 0.5% instead of 5%. A related example is investments in climate change mitigations to cope with potential climate change related extreme events. Definitely, a rate of 3.5%, as often used in integrated assessment models [29], can easily illustrate that climate change does not matter. A shock testing of these models reduces even further their internal stopping time horizon.

Example 4.2. (Darwinian fitness). Ramsey [26] had introduced discounting, first of all, as a mathematical device ensuring the convergence of infinite horizon cumulative values. Its various explanations supported by empirical studies were proposed afterwards suggesting that humans and animals place less weights on the future than on the present [28]. A reason is that future rewards run more risk of disappearing. Hence, they should be discounted, where the discount rate is the hazard rate. For example, evidence from selection experiments indicates the existence of a trade-off between short-term and long-term fertility, i.e., the existence of life-history strategy that discounts the future. In other words, natural selection puts a premium on immediate reproductivity. Accordingly, an animal can be treated as a rational optimizer maximizing its Darwinian fitness, that can be taken to be equivalent to maximizing the expected number of offsprings. In a simple case, fitness is defined [28] then as integral $F = \int_0^\infty m(t)s(t)dt$, where $m(t)$ is the expected rate of reproductive output at age t if the animal survives to that age, and $s(t)dt$ is the probability of surviving to age t . It is highly unlikely that an animal is able to learn discount factors (probability density $s(t)$) in order to maximize the Darwinian fitness. The equivalent distribution free stopping time criterion requires observations of only lifetime intervals τ , which can be easily used for adaptive adjustments of life-history strategies.

4.3 Time Declining Discount Rates

This section extends Proposition 4.1 to general time declining discount rates. It also shows that a time declining discount rate can be associated even with a set of mutually exclusive geometrically distributed extreme (stopping time) events. This rate is determined in a sense by the least probable event.

Let us consider now a stream of random variables (r.v.) v_0, v_1, \dots affected by a set of random events including potential catastrophic events. Formally, we can think of v_t as a function $v_t(\omega)$ defined on a probability space $\{\Omega, P\}$ with the set Ω of related random events and the probability measure P on Ω . We assume that v_t does not depend on the “future”, i.e., we assume that $\{\Omega, P\}$ is adapted to a sequence of increasing σ -algebras $A_0 \subseteq A_1 \subseteq \dots$ (subsets of events from Ω , which occur before $t = 0, 1, \dots$), such that v_t is measurable (defined on) w.r.t. A_t . In what follows, all random variables are assumed to be defined on $\{\Omega, P\}$.

Let $\sigma_{k,t} = \sigma(v_k, \dots, v_t)$ be the σ -algebra generated by v_k, \dots, v_t . Consider a stopping time τ , which we define as a r. v. $\tau \in \{0, 1, \dots\}$, such that event, $\{\tau \leq t\}, t = 0, 1, \dots$ does not depend on values v_{t+1}, v_{t+2}, \dots , i.e., $\sigma_{t+1, \infty}$.

Proposition 4.2. *Consider a discounted sum $\sum_{t=0}^{\infty} d_t V_t, d_t = (1+r_t)^{-t}$, where r_t is an increasing positive sequence, $V_t = E v_t$. Then there is a stopping time τ such that $P[\tau \geq t] = d_t$ and*

$$\sum_{t=0}^{\infty} d_t V_t = \sum_{t=0}^{\infty} P[\tau \geq t] E v_t = E \sum_{t=0}^{\tau} v_t. \quad (4.3)$$

Conversely, let $E | v_t |$ is uniformly bounded. Then, for any stopping time τ

$$E \sum_{t=0}^{\tau} v_t = \sum_{t=0}^{\infty} d_t V_t, d_t = P[\tau \geq t],$$

where V_t is conditional expectation:

$$V_t = E[v_t | \tau \geq t]$$

Proof. Consider such any r.v. $\tau, \tau \in \{0, 1, \dots\}$ that $\{\tau \leq t\}$ does not depend on values v_0, \dots, v_{t-1} and $P[\tau = t] = d_t - d_{t+1}, t = 0, 1, 2, \dots$. Clearly, $P[\tau \geq 0] = d_0 - d_1 + d_1 - d_2 + \dots = d_0 = 1, P[\tau \geq t] = d_t$ and

$$\sum_{t=0}^{\infty} d_t V_t = \sum_{t=0}^{\infty} P[\tau \geq t] V_t.$$

Let now $f_t := \sum_{k=0}^t v_k$. From the rearrangement known as the Kolmogorov-Prohorov's theorem it follows that

$$\begin{aligned} E f_{\tau} &= \sum_{t=0}^{\infty} E[f_t; \tau = t] = \sum_{t=0}^{\infty} \sum_{k=0}^t E[v_k; \tau = t] = \sum_{k=0}^{\infty} E[v_k; \tau \geq k] \\ &= \sum_{k=0}^{\infty} P[\tau \geq k] V_k. \end{aligned}$$

where $V_k = E[v_k \mid \tau \geq k]$ and $E[v_t; A]$, denotes unconditional expectation $E[v_t I_A]$, I_A is the indicator function of event A . The last assertion follows from the identity $\{\tau \geq t\} = \{\tau > t - 1\}$, i.e., from the independence of $\{\tau \geq t\}$ on $\sigma_{t,\infty}$. The change in the order of sums is possible due to the uniform boundness of $E \mid v_t \mid$.

Corollary 4.1. *If v_0, v_1, \dots are independent r.v. or $\{\tau \geq t\}, t = 0, 1, 2, \dots$, does not depend on v_0, v_1, \dots, v_{t-1} , then V_t in both cases of Proposition 4.2 is unconditional expectation $V_t = E v_t$. If v_0, v_1, \dots are independent identically distributed r.v., then the Wald's identity follows from Proposition 4.1:*

$$E \sum_{t=0}^{\tau} v_t = E v_0 E \tau.$$

Proof. It follows from the following rearrangements:

$$\sum_{t=0}^{\infty} P[\tau \geq t] = \sum_{k=0}^{\infty} \sum_{t=k}^{\infty} P[\tau = t] = \sum_{t=0}^{\infty} t P[\tau = t] = E \tau.$$

Example 4.3. (Expected catastrophic losses). Assume that a catastrophic event may occur at $t = 0, 1, 2, \dots$ with probability p . It is usually defined as $(1/p)$ -year event, say a 100-year flood. Define τ as the arrival time of the first catastrophe and let $v_t = 0, 0 \leq t \leq \tau - 1, v_\tau = L_\tau$, where L_τ is conditional expected losses given that the event occurs at τ . Since $l_t \neq 0$ only for, $t = \tau$, then the expected (unconditional) losses at τ are:

$$E v_\tau = p L_0 + q p L_1 + q^2 p L_2 + \dots = \sum_{t=0}^{\infty} q^t V_t = \sum_{t=0}^{\infty} P[\tau \geq t] V_t,$$

where $V_t = p L_t$.

The next proposition shows that a set of even geometrically distributed events can induce discounting with time declining discount rates. Let us assume that there is a set of mutually exclusive events (see also Sect.4.4) of "magnitude" $i = 1, \dots, n$. The probability of scenario i is θ_i , $\sum_{i=1}^n \theta_i = 1$ and, conditional on this scenario, the event i occurs for the first time at τ_i with the probability $P[\tau_i = t] = p_i q_i^t, q_i = 1 - p_i, t = 0, 1, \dots$. Thus, the occurrence of events at t is characterized by a mixed geometric distribution $\sum_{i=1}^n \theta_i p_i q_i^t$. Let τ be the arrival time of a first event. Then $d_t = P[\tau \geq t] = \sum_{i=1}^n \theta_i P[\tau_i \geq t]$. Since $P(\tau_i \geq t) = p_i q_i^t + p_i q_i^{t+1} + \dots = q_i^t$, then evaluation (4.1) takes the form

$$V = \sum_{t=0}^{\infty} d_t V_t, d_t = \sum_{i=1}^n \theta_i q_i^t. \quad (4.4)$$

This equation essentially modifies the standard geometric discounting. Nevertheless, the induced discount factors d_t for large t tend to be defined by the smallest

discount rate of the least probable event. The following proposition is similar to the conclusion in [31].

Proposition 4.3. *Discount factor $d_t = \sum_{i=1}^n \theta_i q_i^t$ in (4.4) is determined for $t \rightarrow \infty$ by the standard geometric discount factor $q_{i^*}^t$ associated with the least probable event i^* ,*

$$p_{i^*} = \min_i p_i : d_t / q_{i^*}^t \rightarrow \theta_{i^*} \text{ for } t \rightarrow \infty.$$

Proof. $d_t = q_{i^*}^t \sum_{i=1}^n \theta_i \chi_i(t)$, where $\chi_i(t) = (q_i / q_{i^*})^t$. From $p_{i^*} < p_i$, $p_i = 1 - q_i$, it follows that $\chi_i(t) \rightarrow 0, t \rightarrow \infty$, for $i \neq i^*$ and $\chi_{i^*}(t) = 1$. Hence, $d_t / q_{i^*}^t \rightarrow \theta_{i^*}$ for $t \rightarrow \infty$.

Remark 4.4. (Finite time horizon T). Propositions 4.1, 4.2, 4.3 hold true also for a finite time horizon $T < \infty$ after substituting probabilities $P[\tau = t]$, $P[\tau \geq t]$ by conditional probabilities $P[\tau = t \mid \tau \leq T]$ and $P[\tau \geq t \mid \tau \leq T]$.

Remark 4.5. (Distribution free approach). Propositions 4.1, 4.2 provide two alternative approaches for discounting: standard discounted criterion of the left-hand side of (4.1), (4.2) with an exogenous discounting, or undiscounted criterion of the right hand side with τ defined by random arrival time of stopping time events. Proposition 4.3 shows that the corresponding induced discounting $d_t = P[\tau \geq t]$ can be a complex implicit function of spatio-temporal patterns of events. The next section illustrates, that τ may depend also on various decisions. All these make it rather difficult to evaluate exact risk profiles $P[\tau \geq t]$ and exogenous discount factors d_t . Therefore, this would require the use of the distribution-free random stopping time criterion and STO methods rather than the standard distribution-based discounted criterion and deterministic optimization methods.

4.4 Endogenous Discounting

This section summarizes typical motivations for developing spatio-temporal catastrophic risk management models with rather natural versions of the stopping time concepts. A typical model may include often the following loop and the potential for positive feedbacks, branching and disequilibrium:

1. Stopping time induces discounting in the form of dynamic risk profiles $d_t = P[\tau \geq t]$.
2. The discounting affects optimal mitigation efforts.
3. Mitigation efforts affect the stopping time τ , risk profiles $P[\tau \geq t]$ and the discounting d_t (return to point 1).

This means that the stopping time criterion induces endogenous spatio-temporal endogenous discounting.

Example 4.4. (Evaluation of a Flood Management Program). Consider a simple version of the catastrophic flood management model developed for the Upper Tisza

river region [9]. The spatio-temporal structure of this model was motivated by the following reasons.

Throughout the world, the losses from floods and other natural disasters are mainly absorbed by the immediate victims and their governments [13]. The insurance industry and its premium payers also absorb a portion of catastrophic losses, but even in the wealthy countries this share is relatively small. With increasing losses from floods, governments are concerned with escalating costs for flood prevention, flood response, compensation to victims, and public infrastructure repair. As a new policy, many officials would like to increase the responsibility of individuals and local governments for flood risks and losses [25], but this is possible only through location-specific analysis of risk exposures and potential losses, the mutual interdependencies of these losses, and the sensitivities of the losses to new risk management strategies.

This is a methodologically challenging task requiring at least the development of spatio-temporal catastrophe models [5, 8, 9, 30]. Although rich data usually exist on aggregate levels, the sufficient location specific data are not available, especially data relevant to new policies. Moreover, catastrophes affect large territories and communities producing mutually dependent losses with analytically intractable multidimensional probability distributions dependent also on various decisions. This critically distinguishes the arising problems from a standard risk management situations, e.g., the well-known asset-liability management. The standard methods, in particular, the existing extreme event theory, are not applicable to rational management of catastrophic risks. The new GIS-based catastrophe models [9,30] are needed to simulate the occurrence of potential extreme events and the samples of mutually dependent catastrophic losses for which no or very few historic observations exist.

In general, a catastrophe model represents the study region by grids, e.g., a relatively small pilot Upper Tisza region is represented by $1,500 \times 1,500$ grids [9]. Depending on the purpose of the study, these grids are aggregated into a much smaller number of cells (locations, compartments) $j = 1, 2, \dots, m$. These cells may correspond to a collection of households at a certain site, a collection of grids with similar land-use characteristics, or an administrative district or grid with a segment of gas pipeline. The choice of cells provides a desirable representation of losses. Accordingly, cells are characterized by their content, in general, not necessarily in monetary units. Values can be measured in real terms, without using an aggregate dollar value. The content of cells is characterized by the vulnerability curves calculating random damages to crops, buildings, infrastructure, etc., under a simulated catastrophic scenario.

Catastrophic floods which are simulated by the catastrophe model, affect at random different cells and produce mutually dependent random losses L_j^t , $j = 1, \dots, m$, from a catastrophic event at time t . These losses can be modified by various decisions. Some of the decisions reduce losses, say a dike, whereas others spread them on a regional, national, and international level, e.g., insurance contracts. If $x = (x_1, x_2, \dots, x_n)$ is the vector of the decision variables, then L_j^t is a random function $L_j^t(x)$.

Flood occurrences in the region are modeled according to specified probabilistic scenarios of catastrophic rainfalls and the reliability of dikes. There are three dikes allocated along the region's river branch. Each of them may break after the occurrence at a random time of a 100-, 150-, 500-, and 1,000-year rainfall characterized by the so-called up-stream discharge curves calculating the amount of discharged water to the river branch per unit of time. In fact, the discharge curves upscale the information about complex rainfall and run-off processes affected by land-use and land-transformation policies. This brings considerable uncertainty in the definition of a $1/p$ -year flood, $p = 1/100, 1/150, 1/500, 1/1,000$. Therefore, a 100-year discharge curve may represent, in fact, a set of floods with different frequencies p , say, $1/150 \leq p \leq 1/100$. In addition to the interval, the uncertainty about p can be given by a prior distribution. Therefore, a single discharge curve, in general, corresponds to a set of $1/p$ -year floods, where p is characterized by a prior probability distribution. For example, it can be characterized by a finite number of probabilistic scenarios p_1, \dots, p_n with prior probabilities $\theta_1, \dots, \theta_n$ as in Proposition 4.3.

The stopping time can be defined differently, depending on the purpose of the policy analysis. A catastrophic flood in our example occurs due to the break of one of the three dikes. These events are considered as mutually exclusive events, since the break of a dike in the pilot region releases the "pressure" on other dikes. Therefore, the stopping time τ can be defined as the first time moment of a dike break. In this case, the probability or induced discount factor $d_t = P[\tau \geq t]$ is an implicit function of t , probabilities $\theta_i, p_i, i = 1, \dots, n$, and the probability of a dike break. The situation is complicated further by the deterioration of dikes in time and/or by inappropriate maintenance of the flood protection system (see also Sect. 4.6), e.g., modifications to the dikes, the removal of some of them, and building new retention areas and reservoirs. Besides these structural decisions, the stopping time τ can be affected by other decisions, e.g., land use policies. Accordingly, depending on goals, the definition of stopping time τ can be further modified. For example, let us assume that the region [10] participates in the flood management program through payments to a mutual catastrophe fund, which has to support a flood protection system and compensates losses to victims. To enforce the participation in the program, the government provides only partial coverages of losses. The stability of this program critically depends on the insolvency of the fund that may require a new definition of τ . Let β be a fixed investment rate enabling the support of the system of dikes on a certain safety level and ξ be a random time of a first catastrophic flood. Denote by L_j^ξ random losses at location $j, j = \overline{1, m}$, at time $t = \xi$ and by π_j the premium rate paid by location j to the mutual catastrophe fund. Then, its accumulated risk reserve at time ξ together with a fixed partial compensation of losses $\chi \sum_j L_j^\xi$ by the government is $R_\xi = \xi \sum_j \pi_j + \chi \sum_j L_j^\xi - \sum_j \varphi_j L_j^\xi - \beta \xi$, where $0 \leq \varphi_j \leq 1$, is the portion of losses compensated by the fund at location j . Let us also assume that the functioning of the flood management program is considered as a long-term activity assuming that growth and aging processes compensate each other. Then, the insolvency of the fund is associated with the event:

$$\xi \sum_j \pi_j + \chi \sum_j L_j^\xi - \sum_j \varphi_j L_j^\xi - \beta \xi < 0. \quad (4.5)$$

Inequality (4.5) defines extreme random events affected by various feasible decisions x including components $(\pi_j, \varphi_j, \chi, b_j, \beta, j = \overline{1, m})$. The likelihood of event (4.5) determines the vulnerability of the program. It is more natural now to define the stopping time τ as the first time when event (4.5) occurs. In this case τ would depend on all components of vector x and the induced discounting would focus on time horizons associated with the occurrence of the event (4.5).

4.5 Dynamic Risk Profiles and CVaR Risk Measure

Example 4.5 illustrates that the probability distributions $P[\tau \geq t]$, $t = 0, 1, \dots$, itself represent key safety characteristics of catastrophic risk management programs. Induced discounting $d_t = P[\tau \geq t]$ then “controls” these risk profiles implicitly through their contributions to discounted goals of programs. Another possibility as this section shows is to impose explicitly safety constraints of the type $P[\tau \geq t] \geq \gamma_t$, for some safety levels γ_t , $t = 0, 1, \dots$. In this case resulting robust strategies would directly control the safety constraints.

Example 4.5. (Safety constraints). The occurrence of disasters is often associated with the likelihood of some processes abruptly passing “vital” thresholds. This is a typical situation for insurance, where the risk process is defined similar to (4.5) by flows of premiums and claims whereas thresholds are defined by insolvency constraints. A similar situation arises in the control of environmental targets and in the design of disaster management programs [5, 8, 9]. Assume that there is a random process R_t and the threshold is defined by a random ρ_t . In spatial modeling, R_t and ρ_t can be large-dimensional vectors reflecting the overall situation in different locations of a region. Let us define the stopping time τ as the first time moment t when R_t is below ρ_t . By introducing appropriate risk management decisions x it is often possible to affect R_t and ρ_t in order to ensure the safety constraints $P[R_t \geq \rho_t] \geq \gamma$, for some safety level γ , or γ_t , $t = 0, 1, 2, \dots$.

The use of this type safety constraints is a rather standard approach for coping with risks in the insurance, finance, and nuclear industries. For example, the safety regulations of nuclear plants assume that the violation of safety constraints may occur only once in 10^7 years, i.e., $\gamma = 1 - 10^{-7}$. It is remarkable that the use of stopping time criterion as in the right-hand side of (4.2) has strong connections with the dynamic safety constraints and dynamic versions of static CVaR risk measures [27]. Let us illustrate this by using the simplest version of climate change stabilization models discussed in [23].

Assume that $R_t = \sum_{k=0}^t x_k$, where decision variables

$$x_k \geq 0, k = 0, 1, \dots, t, \quad t \leq T < \infty.$$

We can consider x_k to be a CO₂ emission reduction at the beginning of period k . At time t the target value on total emission reduction R_t in period t is given as a

random variable ρ_t . It is assumed that the exact value of ρ_t may be revealed at a random period τ , $P[\tau \geq t] = d_t$. The decision path $x = (x_0, x_1, \dots, x_T)$ has to be chosen ex-ante in period $t = 0$ to mitigate climate change impacts associated with the case $R_\tau < \rho_\tau$. Consider the loss function associated with emission mitigation strategy x and given τ :

$$V(x) = E \sum_{t=0}^{\tau} [c_t x_t + b_t \max \{0, \rho_t - R_t\} I_{t=\tau}], \tag{4.6}$$

where deterministic coefficients c_t can be viewed as marginal costs, and b_t as risk factors.

This can be written (Example 4.3) as

$$V(x) = \sum_{t=0}^T d_t \left[c_t x_t + b_t \operatorname{E} \max \left\{ 0, \rho_t - \sum_{k=0}^t x_k \right\} \right].$$

Assume that $V(x)$ is a continuously differentiable function, e.g., a component of random vector $\rho = (\rho_0, \rho_1, \dots, \rho_T)$ has a continuous density function. Also, assume for now that there exists a positive optimal solution $x^* = (x_0^*, x_1^*, \dots, x_T^*)$, $x_t^* > 0$, minimizing $V(x)$ subject to $x_t \geq 0, t = 0, 1, \dots, T$. Then, from the optimality condition for stochastic minimax problems (see discussions in [6], p. 16) it follows that for $x = x^*$,

$$V_{x_t} = c_t - \sum_{k=t}^T b_k P \left[\sum_{s=0}^k x_s \leq \rho_k \right] = 0, t = 0, 1, \dots, T.$$

From this it follows sequentially for $t = T, T - 1, \dots, 0$,

$$P \left[\sum_{k=0}^T x_k \leq \rho_T \right] = c_T / b_T, P \left[\sum_{k=0}^t x_k \leq \rho_t \right] = (c_t - c_{t+1}) / b_t, t = 0, 1, \dots, T-1. \tag{4.7}$$

Since $\operatorname{E} \max \{0, \rho_t - R_t\} = E \rho_t I_{\rho_t \geq R_t} - R_t P[\rho_t \geq R_t]$, then from (4.7) it follows that $V(x^*) = E p b_\tau I_{\rho_\tau \geq R_\tau}$, which can be viewed as a dynamic CVaR (Conditional-Value-at-Risk) risk measure. Equations (4.7) can be used to control dynamic risk profiles, say, profiles with a given safety level γ as in Example 4.5:

$$1 - \gamma = c_T / b_T = (c_t - c_{t+1}) / b_t, t = 0, 1, \dots, T - 1,$$

by appropriate choice of risk factors b_t similar to stationary CVaR risk measures. In this case the minimization of (4.6) controls safety constraints (4.7) with given safety level γ , i.e.,

$$P \left[\sum_{k=0}^t x_k \leq \rho_k \right] = 1 - \gamma, \quad t = 0, 1, \dots, T. \tag{4.8}$$

This is a remarkable result, since the safety constraints, as a rule, are non-convex and even discontinuous, whereas the minimization of function (4.6) is often a convex problem for important practical cases.

Equations (4.7) are derived so far from the existence of the positive optimal solution x^* . The following proposition clarifies this assumption.

Proposition 4.4. *The existence of positive optimal solution follows from $c_T/d_T < 1$, $(c_t - c_{t+1})/d_t < 1$, $t = 0, 1, \dots, T-1$, and the monotonicity of quantiles $\beta_t, \beta_0 < \beta_1 < \dots < \beta_T$ defined by equations*

$$P[\beta_T \leq \rho_T] = c_T/d_T, P[\beta_t \leq \rho_t] = (c_t - c_{t+1})/d_t, t = 0, 1, \dots, T-1$$

Proof. Indeed, the first requirement guarantees that $x_0^* > 0$, $\sum_{k=0}^t x_k^* > 0$, $t = 1, 2, \dots, T$. From the second requirement it follows that $x_0^* + x_1^* > x_0^*$, i.e., $x_1^* > 0$, and so on.

Let us note that in general cases outlined in Example 4.5, process R_t is given by stochastic equations $R_{t+1} - R_t = g(t, x_t)$, $t = 0, 1, \dots, T-1$, where $g(t, x_t)$ is a random function. In this case (4.7), (4.8) would have a form of conditional expectation rather than quantiles. It is even easy to see for $g(t, x_t) = a_t x_t$, where a_t are random variables. In rather general cases a minimization problem (4.6) can be solved by distribution-free stochastic optimization methods proposed in [5, 7–9], i.e., methods which don't use (in general) exact probability distributions.

Remark 4.6. (Robust decision). The stopping time τ in model (4.6) is not associated with the violation of safety constraint (4.8). In catastrophic risk management the model (4.6) is usually considered as an auxiliary submodel. For example, if random ρ_t are affected by a set of decisions y with a cost function $F(y)$, then the minimization of function $V(x) + F(y)$ yields robust decision minimizing total costs under safety constraints (4.8) and a dynamic version of the CVaR risk measure.

4.6 Intertemporal Inconsistency

The time consistency of discounting means that the evaluation of an investment project today ($t = 0$), will have the same discount factor as the evaluation of the same project after any time interval $[0, T]$ in the future. In other words, despite delayed implementation of the project we always found ourselves in the same environment. Only geometric or exponential discounting, $d_t = d^t = e^{\ln(d)t} = e^{-\lambda t}$, where $\lambda = -\ln(d)$, defines a homogeneous time consistent preference:

$$\sum_{t=0}^{\infty} d^t V_t = V_0 + dV_1 + \dots + d^{T-1}V_{T-1} + d^T[V_T + dV_{T+1} + \dots]$$

This is also connected with the geometric probability distribution of the discount-related stopping time τ in (4.2): if $P[\tau \geq t] = d^t, 0 < d < 1$, then

$$P[\tau = t] = d^t - d^{t+1} = (1 - d)d^t, t = 0, 1, \dots$$

In other words, the consistency is the direct consequence of the well-known “memoryless” feature of geometric and exponential probability distributions: for any $t \geq 0, s \geq 0$,

$$P[\tau = t + s \mid \tau \geq t] = d^{t+s}(1 - d)/d^t = d^s(1 - d)$$

Hence, independently of waiting time t , the probability of the stopping time occurrence at $t + s$ is the same as at the initial time moment $t = 0$.

For other discount factors with time-dependent rates, their time inconsistency requires appropriate adjustments of discount factors for projects undertaken later rather than earlier. The misperception of this inconsistency may provoke increasing vulnerability and catastrophic losses. Let us consider typical scenarios of such developments. Section 4.4 shows that the adequate perception of proper discounting is a challenging task requiring models that allow the explicit evaluation of related risk profiles. This section, in fact, illustrates that the design of such models has to be considered as a key mitigation measure to cope with increasing vulnerability.

A number of authors distinguish between various types of so-called “imperfect altruism” resulting in the lack of social commitment to mitigate risks. For example, there were alluded definitions of a naive, a sophisticated and a committed (ideal) society. The main differences between these three societies and how they provoke catastrophes are summarized in [11] by using a simplified flood management model outlined in Sect. 4.4. This model has the fixed 100-year horizon T in which three societies, the naive, the sophisticated, and the committed, live and plan for coping with the catastrophic losses that may occur due to break of a dyke from 150-year flood with time consistent geometric probability distribution. They are able to mitigate the reliability of dikes and losses by paying fair premiums to the catastrophe fund. But, depending on their perception of risk profiles or induced discounting, the results are dramatically different.

The current generation of *The Naive Society* is aware of a possible catastrophe. It maximizes the (identical for all generations) value function taking into account the potential need to save for paying premiums. Unfortunately, it has a misleading view on the catastrophe, namely, if the catastrophe has not occurred in the later generation the society believes that it will not occur within the current generation with the same probability. Thus, it relies on geometric probability distribution and fails to take into account the time inconsistency induced by increasing the probability of a dike break due to aging processes. Therefore, the first generation of the society postpones the implementation of decisions, i.e., the naive society puts also its preferences on consumption as the first priority consuming at a higher rate than it actually plans.

For the next generation the time is shifted forward by 20 years, and the second generation, similar to the first, plans but does not implement saving actions essential

for the catastrophe fund to function. The risk profiles, time preferences, premiums, and the actions are not adjusted towards the real escalating risks. In a similar way, behave the next generations. The plans are never implemented and the view on a catastrophe remains time invariant despite dramatic increase of risk.

The Sophisticated Society implies a correct understanding of the time-inconsistent discounting induced by the deteriorating system of dikes. But this society, similar to the naive planners, also evaluates present consumption to be much higher than the future one. This leads to postponing the decisions. Due to these delays, the risk burden is increasingly shifted to the next generation, calculated premiums become higher and higher. If a catastrophe occurs, this society will also be not prepared to cope with losses as catastrophe management is not functioning.

The “pathologies” of these societies can be explained by their misperception of risks, and, the lack of committed actions.

The Committed Society is similar to that of the sophisticated society. In contrast though, this society is able to implement decisions because its calculations demonstrate that the delays in actions may dramatically affect individuals and the growth of societies as a whole. Individuals could be better off if their consumption options were limited and their choices constrained by anticipating risks. As a direct consequence of the committed actions, the premiums that the society pays for coping with catastrophes in 100 years time are much lower than those of the sophisticated society.

4.7 Concluding Remarks

The proposed new approach to discounting is based on undiscounted stopping-time criterion which is equivalent to the standard discounted criterion in the case of market-related discount factors. In general, the stopping time criterion induces the discounting that depends on spatio-temporal patterns of catastrophes and various relevant decisions. More formally, this paper demonstrates that discount factors $d_t, t = 0, 1, \dots$ can be associated with the occurrence of an extreme (“killing”) “stopping time” event at random time τ with probability $P[\tau \geq t] = d_t$. Consequently, the infinite discounted sum $\sum_{t=0}^{\infty} d_t V_t, V_t = E v_t$, is replaced by the undiscounted expectation $E \sum_{t=0}^{\tau} v_t$ within the finite interval $[0, \tau]$. The use of the stopping time criterion $E \sum_{t=0}^{\tau} V_t$ induces the standard discounting in the case when τ is associated with the lifetime of market products. In dealing with catastrophic risks, the stopping time τ can be associated with the arrival time of potential catastrophic events. The use of random criterion $\sum_{t=0}^{\tau} v_t$ allows to address the variability of valuations even in the case of deterministic flows V_0, V_1, \dots . In this case, it is often important to substitute the expected value of random sum $\sum_{t=0}^{\tau} v_t$ by its quantiles. Mitigation efforts affect the occurrence of extreme events and, thus, they affect discounting, which in turn affects mitigations. This endogeneity of discounting restricts exact evaluations of d_t and the consequent use of deterministic methods and it calls for specific stochastic optimization methods.

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Chapter 5

Cost Effective and Environmentally Safe Emission Trading Under Uncertainty

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Abstract The aim of this paper is to analyze robust cost-effective and environmentally safe carbon emission trading schemes under uncertainties of emissions and costs, and asymmetric information of participants. The proposed model allows to control explicitly the safety of Kyoto (or other) targets by taking long-term perspectives on emission trading. The dynamics of this scheme is driven by bilateral trades with different endogenous disequilibrium prices between mutually beneficial trades, but finally the system converges to cost-effective and environmentally safe global equilibrium. The safety constraints work as a discounting mechanism that discounts the reported emissions to detectable undershooting levels. This, in turn, provides incentives for participants to reduce uncertainties. The model shows that uncertainties and short term market perspectives may easily prevent price-based trading to be environmentally safe and cost-effective scheme. The desirable equilibrium emerges only under proper price-formation mechanisms. The role of the proposed computerized multi-agent trading system is central for dealing with long-term perspectives, irreversibility and lock-in equilibriums of trades. This system can be viewed as a device for decentralized collective regulation of trades based on unified approaches to modeling of uncertainty, calculation of costs and trading rules.

5.1 Introduction

The public property of large scale pollution makes it impossible to organize complete environmental markets with private demand for and private supply of pollution control [2, 3, 7, 15, 18, 20, 22]. Yet, the idea of carbon trading markets is becoming increasingly popular for global climate change control. At the same time, the existence of various exogenous and endogenous inherent uncertainties raises serious concerns regarding the ability of carbon trading markets to

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fulfill the main purpose of the climate change control without creating world-wide irreversible socio-economic and environmental disruptions. Definitely, that interests of profit oriented markets may contradict the main concerns of the Kyoto agreements [29].

Considered in this paper is bilateral exchange of emission rights. It is assumed that parties with high emission reduction costs buy emissions from parties with low emission reduction costs within prescribed targets. In other words, parties can be engaged in a mutually beneficial bilateral emission exchange process [8] driven by cost minimizing and environmentally safe bilateral trades without the need for a market. This approach is close to important ideas on decentralized non-monetary exchange [21, 28].

In contrast, carbon trading markets, which become increasingly popular in recent years, are more similar to stock markets. Parties hold a number of permits to emit a specific amount of emissions. Parties that cannot to keep their emissions at the given level (called cap) must buy permits on the market at a prevailing market price.

Thus, in the bilateral emission trading scheme the exchange of emission rights is driven by the abatement costs and safety constraints, while in the carbon trading markets the exchange of emissions is driven by prevailing market prices. Such price signals with potential bubbles created by speculators may have no connections with minimization of abatement costs and achieving environmental safety constraints.

There are two principle approaches to control pollution: centralized cooperative command-and-control methods and decentralized market simulating schemes. If there was a social planner (central agency or regulator) fully informed about emissions and abatement cost functions of all parties, the primal problem of finding emission levels that meet given environmental standards in a cost-effective way would be a straightforward task. This could be done by dealing with nonconvex cost functions typically encountered in long-term evaluations involving new technologies with increasing returns. However, without such a planner, the primal model has to be solved in a decentralized manner.

The aim of this paper is to develop an integrated approach for designing cost-effective and environmentally safe decentralized emission trading schemes robust with respect to uncertainties of emissions, costs and asymmetric information of parties. The bilateral emission trading scheme of Sect. 5.5 corresponds to a decentralized solution of the primal model, whereas schemes of Sect. 5.7 simulate decentralized price-based market's solutions. The cost-effectiveness and environmental safety of latter solutions critically depend on proper price signals, which usually reflect instantaneous market situations rather than long-term costs and environmental constraints of the dual model. The complexity of the primal model is a vital issue for the existence of proper prices.

There is a number of uncertainties, affecting outcomes of examined model. First of all, emissions of Green-House Gases (GHGs) are not directly observable. A comprehensive discussion of related issues can be found in the volume by [1], and in [12]. In general, emissions can be estimated with information on the GHG-emitting activities by applying specific conversion factors and from atmospheric measurements using inversion models. The accuracy of the reported emissions depends on

the quality of the monitoring system in each specific country and on the accuracy of the conversion factors used [26]. As emissions of GHGs cannot be observed perfectly, the uncertainty can be misused by concealing unreported emissions. A central issue becomes a trade-off between reductions of emissions and uncertainties. For example, carbon prices in the European Union crashed and caused instabilities in late April 2005 after the Czech Republic, Estonia, France, the Netherlands and Sweden reported lower than anticipated emissions [4, 27].

Apart from emissions, another essential uncertainties are those related to the emission amounts and reduction costs. Parties have incentives to keep this information private and the specific costs may remain unknown to the other parties. Besides, they may vary according to unknown market conditions. They are also subject to both industry wide and firm specific shocks.

The novelty of this paper is in integrated analysis of emission trading schemes under various types of natural and human related uncertainties. Section 5.2 illustrates the need for proper treatment of uncertainties by using available historical observations of CO₂ emissions. It shows that the use of uncertainty intervals as practiced by International Panel for Climate Change (IPCC) can leave out of consideration an essential mass of potential emissions. Therefore, Sect. 5.3 introduces a simple and realistic stochastic model allowing to represent both, human related uncertainties and uncertainties associated with the natural variability of emissions. The model focuses on proper representation of potentially controversial experts' judgments and path-dependencies of emissions. As a result, this allows us to introduce safety constraints and undershooting mechanism to control the robustness of emission targets during trading process.

Section 5.4 introduces a basic model allowing to analyze different trading schemes. In particular, the model shows that the trade equilibrium under uncertainty is significantly affected by uncertainty. This emphasizes the need for proposed integrated modeling of uncertainties, safety constraints and emission trading schemes.

The dynamic bilateral trading scheme of Sect. 5.5 can be viewed as a stochastic decomposition procedure. The trade at each step takes place towards minimization of safety-adjusted costs of meeting parties. This generates disequilibrium random prices which are endogenously driven towards the cost-effective and environmentally safe equilibrium price. This section analyses also difficulties involved in setting up such an equilibrium price in monetary trading schemes. Standard market models usually imply (e.g., by an arbitrage free type of assumption) that markets operate under equilibrium prices. Section 5.6 outlines a computerized Multi-Agent Decentralized Trading System dealing with the irreversibility of emission trades. Section 5.7 analyses path-dependencies of myopic trading schemes relying on instantaneous markets situations. It shows that short-term market perspectives preclude achieving desirable long-term emission reduction goals. Section 5.8 concludes and outlines important numerical results. The Appendix provides a proof of the convergence. It also discusses stable core solutions of bilateral emission trading scheme.

5.2 Uncertainties and Trends of Carbon Fluxes

This section illustrates a general need for proper representation of emission uncertainties. Next section addresses these issues in a more specific context of emission trading.

Uncertainties of emissions are often represented by means of intervals. In reality, emissions may have different likelihoods within these intervals, i.e., rather general skewed probability distributions. In this case, the use of uncertainty intervals can leave out of consideration essential patterns of emission changes as in Figs. 5.2 and 5.3.

Figures 5.1, 5.2, and 5.3 illustrate trends and natural variability of net carbon fluxes on the global scale <http://lmacweb.env.uea.ac.uk/lequere/co2/carbon.budget.htm>. The global carbon budget is composed of the fossil fuel emissions, the emissions stemming from land use, the ocean uptake, and the terrestrial uptake estimated as a residual of all the sources minus the ocean uptake and atmosphere increase.

Figures 5.2 and 5.3 show the dynamics of changes in emissions and emissions uncertainties. The histogram in Fig. 5.2 is skewed to the left. In the next study period, Fig. 5.3, the situation changes: more values are concentrated on the right hand side. Between these two periods, the system turns from sink to source of CO₂. Definitely, it is impossible to characterize these changes only by uncertainty intervals.

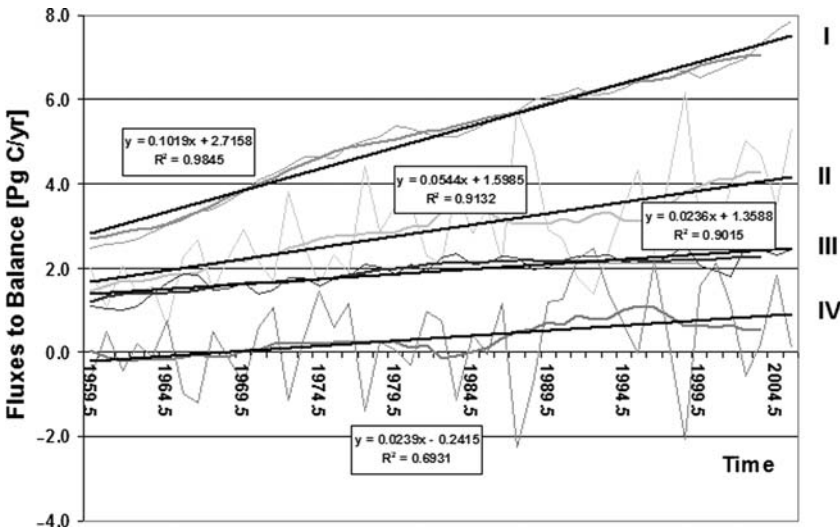


Fig. 5.1 Emission trends: fossil fuel and cement burning (I); CO₂ in the atmosphere (II); mean ocean uptake (III); net terrestrial flux (IV). *Bold lines* correspond to smoothed trajectories of respective fluxes. Regression equations in *boxes* describe linear trends

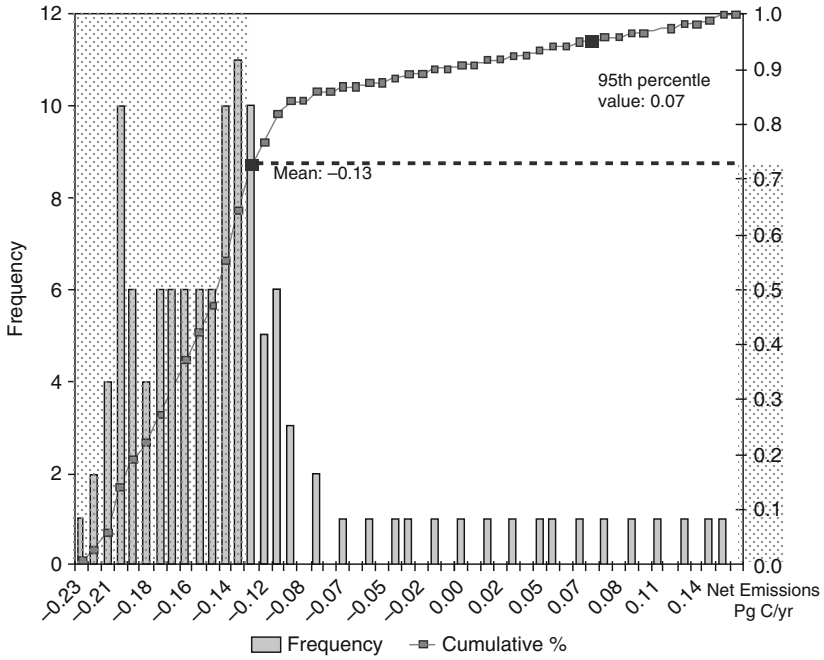


Fig. 5.2 Global CO₂ net terrestrial uptake, 1960–1970

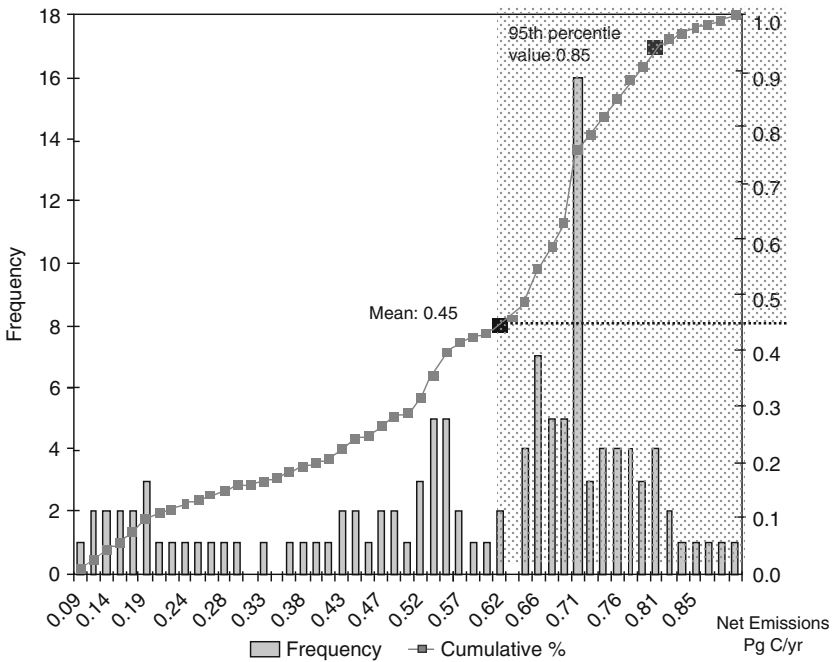


Fig. 5.3 Global CO₂ net terrestrial uptake, 1985–1995

5.3 Detectability of Emission Changes

A simple way to introduce the detectability of emission changes can be based on a straightforward representation of emission trends and uncertainties by equally-sided intervals as in Fig. 5.4 (see [1, 19]). The main idea is illustrated in Fig. 5.4. Let us assume that uncertainty of emission e_1 in the base year t_1 is characterized by equally sided interval $[e_1 - \epsilon, e_1 + \epsilon]$. The uncertainty of reported emission e_2 in the commitment year t_2 ($t_1 < t_2$) is characterized by the same type of interval $[e_2 - \epsilon, e_2 + \epsilon]$. We assume that $e_1 > e_2$, although the case $e_1 < e_2$ is also possible, e.g., as a result of emission trading. The detectability of emission changes requires that the change in net carbon emissions $\Delta e = e_1 - e_2$ at time t_2 is greater than the uncertainty in the reported net carbon emissions at time t_2 .

Under the non-restrictive assumption that the first-order linear approximations for emissions (as in Fig. 5.1), $e(t)$ and uncertainties $\epsilon(t)$ trends are applicable for $t_1 \leq t \leq t_2$, the detection time t^* is defined as the first time moment when net emission change Δe outstrips the uncertainty interval. In a sense, this is a worst-case evaluation. As in Figs. 5.2 and 5.3, considerable mass of real emissions can be concentrated in a much smaller subinterval. For example, the uncertainty interval of random variable with a normal probability distribution is $(-\infty, \infty)$, whereas practically entire probability mass may be concentrated within $[-1, 1]$ interval. Therefore, by using stochastic uncertainty models it is possible to derive with high probability a more optimistic t^* . This is the main issue of stochastic models discussed in [16, 17]. An overview of different approaches can be found in [1]. The goal of the stochastic models is to rank the trading parties by a safety indicator showing the percentage of detectable emission changes within a given time interval.

Let us consider a rather general stochastic model for a representation of a controversial data about uncertainties. We assume that the uncertainty of emissions e_1 ,

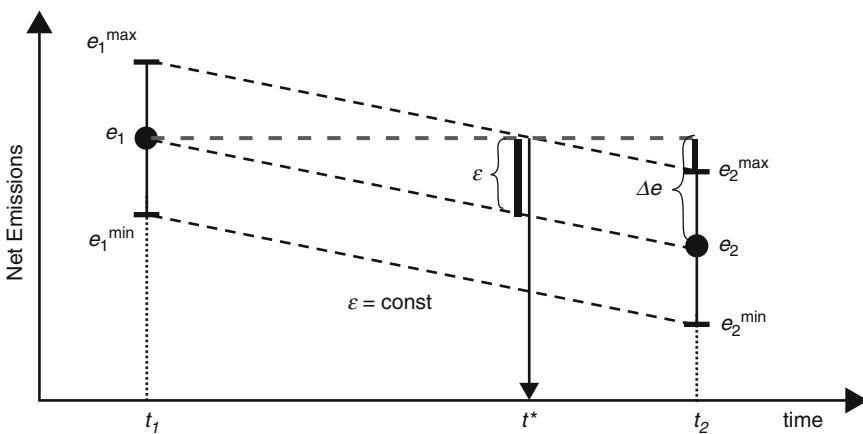


Fig. 5.4 Simplified illustration of detection time t^*

e_2 is characterized by a set of, in general, disconnected intervals. For example, Figs. 5.2 and 5.3 may suggest to represent uncertainty by a number of subintervals characterized by simple, say, uniform (conditional) probability distributions.

The following simple example illustrates the main idea of the model.

Example 5.1 (Controversial experts). Experts judgments are used in situations with the lack or even absence of real observations. Assume that two experts, Ex.1 and Ex.2, characterize the uncertainty of e_1 by overlapping intervals $[1, 8]$, $[5, 10]$. Then overall uncertainty of e_1 can be characterized by intervals $[1, 5]$, $[5, 8]$, $[8, 10]$ with likelihoods $1/4$, $2/4$, $1/4$ derived from the “voting” of experts: $(1, 0)$, $(1, 1)$, $(0, 1)$, i.e., for interval $[1, 5]$ votes only Ex.1, both experts vote for interval $[5, 8]$, and only Ex.2 votes for $[8, 10]$. In general, experts may characterize uncertainty by disconnected intervals. For example, Ex.1 may insist on equally probable intervals $[1, 3]$, $[5, 8]$ conditional on implementation of different technologies. Uncertainty of e_2 can be characterized in a similar manner.

Consider now a general model. For simplicity of notation we omit the index i of parties. Assume that (specific for each party i) uncertain emission e_1 is characterized by intervals $[e_1^{\min}, e_1^1]$, $[e_1^2, e_1^3]$, $[e_1^4, e_1^5]$, \dots , $[e_1^R, e_1^{\max}]$ with probabilities p_r , $\sum_r p_r = 1$. These intervals can be derived from real observations, experts opinions, and scenarios of future developments. In addition, likelihoods of emissions within an interval can be characterized by a conditional on r distribution, say, uniform, normal or the degenerated distribution concentrated in the middle of this interval as in Fig. 5.4. In a similar manner, emissions e_2 are characterized by intervals $[e_2^{\min}, e_2^1]$, $[e_2^2, e_2^3]$, $[e_2^4, e_2^5]$, \dots , $[e_2^L, e_2^{\max}]$ with some conditional on r and l distributions. Path-dependencies between emissions e_1 and e_2 are induced by the following stochastic model. An interval r at the base year t_1 is selected with the probability p_r , emission level e_1 is sampled from the conditional distribution in this interval; an interval l of a trend from r to l is selected with probability q_{rl} , $\sum_l q_{rl} = 1$, and finally, the end point e_2 of the random linear path (e_1, e_2) is sampled from the distribution in interval l conditional on r and l . Let us denote the obtained linear random path by $e(t, \omega)$, $t_1 \leq t \leq t_2$, where ω denotes the pair of points (e_1, e_2) .

Linear paths $e(t, \omega)$ create the uncertainty ranges at t_2 . For example, if $e(t_1, \omega)$ belongs to interval r , then the uncertainty of e_2 is defined on the basis of only feasible transitions from interval r to random intervals l with positive q_{rl} , $q_{rl} > 0$.

The proposed stochastic model allows to introduce path-dependencies of emissions subject to some essential conditions, say, the implementation of new emission reduction technologies or monitoring equipment. This may simplify the detection of emission changes. Exact detection is in general a difficult task because the resolution of all involved uncertainties may be prohibitively costly. Yet, it is possible to define likelihoods of changes. For example, it is possible to find a minimal time t_2 such that emission changes are detected during $[t_1, t_2]$ with a specified likelihood. It is also possible to find the likelihood of the changes within given interval $[t_1, t_2]$ that is used in the next section.

Remark 5.1 (Modifications of model). The proposed stochastic model can be further generalized or simplified subject to available data. Straight lines of linear

emission paths can be theoretically substituted by more general stochastic paths (processes) or scenario trees, although the proposed linear stochastic trends allow simple calculations. If path-dependencies of emissions are not essential, then the model directly deals only with uncertainties of e_2 . In more general situations, uncertainties are also characterized by a set of probability distributions, i.e., there is a set of feasible p_r, q_{rl} . There exist different approaches to deal with arising “uncertainty-of-uncertainty” issues, in particular, the use of non-Bayesian worst-case distributions [6].

Comparative analysis of deterministic and stochastic simple detection models can be found at <http://www.iiasa.ac.at/Research/FOR/unc-prep.html>, [FOR/vt-concept.html](http://www.iiasa.ac.at/Research/FOR/vt-concept.html), and in [10].

5.4 Trade Equilibrium Under Uncertainty

GHG’s control policy as other environmental policies have to be designed in such a way that they are environmentally safe and cost-effective. The models proposed in this section provide a basis for designing rather different decentralized emission trading schemes.

The models reflect the following key features. The participants (countries, companies or other emitting entities) are given a right to emit a specific amount for which they obtain an equivalent number of allowances (emission permits). Such amounts are called the “cap” (Kyoto or other targets). If participants emit more than the corresponding cap reduced by the amount of uncertainty (undershooting level) ensuring that the actual emission does not exceed the cap with a given safety (likelihood) level, they are required to reduce uncertainty or/and to buy additional credits from the parties which emit less than their cap. The transfer of permits is called “trading”. Standard deterministic models belong to a specific class of the proposed models. Since they ignore uncertainty, actual emissions may considerably overshoot allowed targets.

Let us briefly consider a deterministic model with uncertainty intervals proposed in [13, 14], that will be further extended to include stochastic safety constraints. The decision problem of each party can be separated in two interdependent subproblems. Firstly, for a given amount of permits, each party solves individual problem deciding whether to spend resources on abating emissions or investing in uncertainty reduction to satisfy emission targets. This problem does not require the information from any other party. Secondly, the party needs to decide whether or not to exchange permits with other parties. This decision problem involves the cost functions of other parties. In the model this information is private and therefore the methodology of decentralized optimization [8, 10] is required.

For the individual optimization problem, we define the least costs $f_i(y_i)$ for party i to comply with imposed targets for a given amount of permits y_i and the target K_i as the minimum of emission reduction costs $c_i(x_i)$ and costs of uncertainty

reduction (e.g., by investments in monitoring) $d_i(u_i)$:

$$f_i(y_i) = \min_{u_i, x_i} [c_i(x_i) + d_i(u_i)], \quad (5.1)$$

$$x_i + u_i \leq K_i + y_i, x_i \geq 0, u_i \geq 0, \quad (5.2)$$

for all i , where x_i is the reported emissions at source i , u_i is its uncertainty, and y_i is the amount of emission permits acquired by source i (y_i is negative if i is a net supplier of permits). Therefore, constraint (5.2) requires that the reported emission x_i undershoots the target K_i by the level of uncertainty u_i . Similar concept of undershooting is also used in [24]. There are also suggestions [23] to represent uncertainty by a fraction of x_i . Example 5.2 shows that this case can be reduced to the case of additive uncertainty as in (5.2). Let us also note, that the model can be formulated in terms of emission reductions that require only slight changes of terminology.

The second optimization problem with asymmetric information involves finding the permit vector $y = (y_1, \dots, y_n)$ or distribution of permits minimizing unknown total or social cost function

$$F(y) = \sum_{i=1}^n f_i(y_i) \quad (5.3)$$

subject to

$$\sum_{i=1}^n y_i = 0. \quad (5.4)$$

Suppose that the cost functions $c_i(x_i)$ and $d_i(u_i)$ are positive, decreasing, convex in x_i and u_i respectively and continuously differentiable. Therefore, $f_i(y_i)$ is also convex, positive, decreasing and differentiable. Then, from the Lagrangian minimization a trade equilibrium can be defined as the vector $y = (y_1, \dots, y_n)$ satisfying the following equations:

$$f'_i(y_i) = -\lambda, \quad \sum_{i=1}^n y_i = 0. \quad (5.5)$$

The condition (5.5) states that the marginal value of a permit shall in equilibrium be equal to a specific unknown level (price) λ same for all parties. It is clear that at the equilibrium vector y^* the constraints (5.2) will hold with equality, i.e.,

$$f_i(y) = \max_{x_i} [c_i(x_i) + d_i(K_i + y_i - x_i)] = \max_{u_i} [c_i(K_i + y_i - u_i) + d_i(u_i)].$$

Therefore from (5.1), (5.2) it follows that at the equilibrium $y_i = y_i^*$, $\lambda = \lambda^*$, $x_i = x_i^*$, $u_i = u_i^*$:

$$c'_i(x_i) = d'_i(u_i) = -\lambda, \quad \sum_{i=1}^n y_i = 0, \quad (5.6)$$

where (x_i^*, u_i^*) is the solution of the subproblem (5.1), (5.2) for (y^*, λ^*) , $y^* = (y_1^*, \dots, y_n^*)$ satisfying (5.5). This equation states that in the cost-effective and environmentally safe equilibrium, the marginal cost of holding emissions down to x_i^* will be equal to the marginal cost of holding uncertainty down to u_i^* . It shows that the explicit introduction of uncertainty u_i and the safety constraints (5.2) into emission trading schemes may significantly affect the equilibrium and, hence, the design of proper emission trading schemes. In particular, it means that equilibrium market prices λ^* must satisfy (5.5), (5.6). In other words, if λ^* is known and f_i are convex functions, then λ^* decentralizes joint model (5.3), (5.4) into individual solutions of (5.5). For non-convex function $F(y)$, (5.5) are not sufficient to find an equilibrium solution of the model.

However, there is no social planner that knows the cost functions of all parties. Therefore, even in the convex case the optimal value of $F(y)$ and λ^* cannot be resolved by solving (5.5). The scheme of bilateral trade presented in Sect. 5.5 allows to compute the equilibrium x_i^* , u_i^* , λ_i^* without revealing private information on functions f_i .

Remark 5.2 (Long-term perspectives, detectability and undershooting). The basic model can be easily extended to a dynamic version. In this article we do not consider it explicitly. Instead, we introduce below long-term perspectives by explicit treatment of future uncertainties and dynamic trading processes. The environmental constraint (5.2) assumes that the known emissions plus the uncertainty of emissions undershoot the emission target. This corresponds exactly to the detectability concept in Fig. 5.4. Constraint (5.2) discounts, in a sense, the reported emissions to levels undershooting emission targets. As (5.6) show, this provides incentives for the uncertainty reduction.

In the stochastic model of Sect. 5.3 uncertainty of emissions by party i at the commitment year t_2 is characterized by a random variable $e(t_2, \omega_i)$. A reported emission x_i provides additional information that modifies $e(t_2, \omega)$. For example, if it is known for sure that x_i belongs to an interval l_i , then distribution of $e_i(t_2, \omega_i)$ is induced only by feasible transitions from initial intervals r_i to l_i with corresponding probability distributions. We can also say that reported emission x_i transforms $e_i(t_2, \omega_i)$ into a random variable $\epsilon_i(x_i, \omega)$. Therefore, (5.2) of the deterministic model has to be understood now in a probabilistic sense as the following safety constraint. Let us define the uncertainty of reported emission x_i as $\xi_i(x_i, \omega) = \epsilon_i(t_2, x_i, \omega) - x_i$. Then the safety constraint can be written as probabilistic version of the deterministic constraint (5.2):

$$P[x_i + \xi_i(x_i, \omega) \leq K_i + y_i] \geq Q_i, \quad (5.7)$$

for all parties i , where Q_i is a safety level ensuring that the probability of all potential emission paths to x_i satisfying the emission target K_i exceeds Q_i . Thus the interval uncertainty u_i is substituted by a random variable $\xi_i(x_i, \omega)$ dependent, in general, on x_i . In reality, the uncertainty characterized by ξ_i can be reduced by improvements of monitoring systems. Let us introduce the variable u_i to control ξ_i

within the desirable safety level Q_i . If $z_i(x_i)$ is the minimal z such that

$$P[\xi_i(x_i, \omega_i) \leq z] \geq Q_i,$$

then the safety constraint (5.7) can be substituted by the following equivalent constraint

$$x_i + u_i \leq K_i + y_i, u_i \leq z_i(x_i), \quad (5.8)$$

Remark 5.3 (Risk-based undershooting). Equation (5.8) shows that the stochastic model induces risk-based upper bounds on uncertainty intervals. Therefore, it allows, e.g., in cases illustrated by Figs. 5.3 and 5.4 to introduce but risk-based undershooting of emission targets defined by “critical” quantile $z_i(x_i)$, which is less conservative than standard interval uncertainty.

For the simplicity of notation, let us denote now by ω the vector of all uncertain parameters affecting cost functions and emissions, i.e., some components of ω such as market prices affect only cost functions, whereas other components affect emissions. In other words, suppose that all uncertain variables are defined on a probability space with a set of scenarios (events) ω . For random cost functions $c_i(x_i, \omega)$, $d_i(u_i, \omega)$, we can redefine functions $f_i(y_i)$ in (5.1) as

$$f_i(y_i) = \min_{x_i, u_i} E[c_i(x_i, \omega) + d_i(u_i, \omega)], \quad (5.9)$$

where the minimization in (5.9) is subject to constraint (5.8). In this model ex-ante decisions x_i , u_i take a long-term perspective: they have to be optimal against all potential future scenarios ω and threats regulated by safety constraints. Uncertainties of cost functions c_i , d_i may be due to unknown in advance market performance, production shocks, and technological uncertainties.

Example 5.2 (Linear equivalent). Often, $\xi_i(x_i, \omega)$ is represented as $\xi_i(x_i, \omega) = \gamma_i x_i + \epsilon_i$, where $0 \leq \gamma_i \leq 1$, and ϵ_i is a random variable. In particular, uncertainty u_i in (5.2) can be given as $u_i = \gamma_i x_i$. The uncertainty in these cases can be controlled by γ_i in the following manner. Let $\epsilon_i(Q)$ be the minimal z such that $P[\epsilon_i \leq z] \geq Q_i$, e.g., $\epsilon_i(Q) = 0$ for constraints (5.2). Then constraint (5.8) is reduced to linear constraint

$$x_i + u_i \leq K_i - \epsilon_i(Q_i) + y_i, u_i \leq x_i.$$

After solving individual subproblem subject to this constraint, the optimal γ_i can be found as $\gamma = u_i / x_i$.

Remark 5.4 (Nonconvexity). Safety constraints (5.7) are well known in financial applications as the Value-at-Risk indicator. Similar constraints are typical for safety regulation of insurance companies, nuclear power plants, and catastrophic risk management [9]. Unfortunately, due to these constraints $f_i(y_i)$ and $F(x)$ may not be a convex function. In order to ensure convexity and/or robustness of decisions x_i , u_i

under rare extreme events it is possible to modify slightly cost functions c_i , d_i based on the “Conditional Value at Risk” [25] indicators as it is used in [9] and in [6] for general stochastic optimization problems.

Additional nonconvexities of functions $f_i(y_i)$ may be due to increasing returns of cost functions $c_i(\cdot)$, $d_i(\cdot)$, with respect to new emission abatement technologies with increasing returns. Nonconvexities generate the so-called duality gap between solutions of the basic model (5.3), (5.4) and its dual model (see Sect. 5.6) precluding the price-based dual (market) schemes to achieve cost-effective and environmentally safe solutions.

5.5 Dynamic Bilateral Trading Processes

The overall goal of the parties participating in the emission trading is to jointly achieve emission targets by redistributing the emission permits y_i , i.e., to find a vector y that would minimize social costs of all parties (5.3) under safety constraints (5.8), where cost functions $f_i(y_i)$ are defined according to (5.9). It is assumed that a party i knows its expected cost function $f_i(y_i)$, but the expected cost function $F(y)$ is unknown.

The basic feature of the trading scheme is similar to the procedure in [8] for convex function $F(y)$: two parties meet (e.g., picked at random) and, if possible, exchange emission permits in a mutually beneficial way. A new pair is picked and the procedure is repeated. The Appendix provides the proof that this dynamic process will lead the parties to an equilibrium despite the information of each party’s cost is private and $F(y)$ is not necessarily a convex function.

The following simple equations illustrate that the bilateral exchange of emissions is beneficial for both parties. Let $y^k = (y_1^k, \dots, y_n^k)$ be the vector of emission permits after k trades. Consider two parties i and j at step k with permits y_i^k and y_j^k . An exchange of permits between them leads to a new distribution of permits $y^{k+1} = (y_1^{k+1}, \dots, y_n^{k+1})$, $y_l^{k+1} = y_l^k$ for $l \neq i, j$. According to (5.5), if there exist any two parties i and j having different marginal costs on emission reduction $f'_i(y_i^k) \neq f'_j(y_j^k)$, then the permit vector $y^k = (y_1^k, \dots, y_n^k)$ is not cost efficient. Without loss of generality, assume that $f'_i(y_i^k) - f'_j(y_j^k) < 0$. Constraint (5.4) requires that the feasible exchange in permits has to be such that $y_i^{k+1} + y_j^{k+1} = y_i^k + y_j^k$.

If we take $y_i^{k+1} = y_i^k + \Delta_k$ and $y_j^{k+1} = y_j^k - \Delta_k$, $\Delta_k > 0$, then the new feasible distribution of permits reduces the total costs of parties $f_i(y_i^k) + f_j(y_j^k)$ and hence the total cost $F(y^k)$:

$$\begin{aligned} F(y^{k+1}) - F(y^k) &= f_i(y_i^{k+1}) + f_j(y_j^{k+1}) - f_i(y_i^k) - f_j(y_j^k) \\ &= \Delta_k(f'_i(y_i^k) - f'_j(y_j^k)) + o(\Delta_k) < 0, \end{aligned}$$

for a small Δ_k . We also have

$$f_i(y_i^{k+1}) - f_i(y_i^k) < f_j(y_j^k) - f_j(y_j^{k+1}). \quad (5.10)$$

i.e., the new distribution of permits reduces costs of j more than increases cost of i . Hence j is able to compensate i for the increased costs in a mutually beneficial way.

Let us summarize the trade scheme more precisely. We assume that after picking up (say at random) a pair of parties i, j these parties are able to find y_i^{k+1}, y_j^{k+1} minimizing

$$f_i(y_i) + f_j(y_j) \quad (5.11)$$

subject to constraints $y_i + y_j = y_i^k + y_j^k, y_i \geq 0, y_j \geq 0$.

This problem is solved by parties i and j only. For continuously differentiable functions $f_i(y_i), f_j(y_j)$, a party j that decreases emission permit by $\Delta_k > 0$ may negotiate with i such a level Δ_k that equalizes marginal costs, i.e., $f_i'(y_i^k - \Delta_k) = f_j'(y_j^k + \Delta_k) = \lambda_k$, where λ_k can be viewed as an equilibrium price (in general stochastic) at step k . Let us note that price process λ_k is driven endogenously by cost-minimizing decisions of meeting parties, what is fundamentally different from standard models of financial markets with exogenously given price processes.

The sequential bilateral trades can go on as long as there are two parties with different marginal costs. The bilateral exchange of emissions equalizes marginal costs which define an intermediate “local” equilibrium price λ_k . During the process, marginal costs and prices will differ between the sequential trades, but finally the trading system converges to an equilibrium with marginal costs of all parties equal to equilibrium price as in (5.5).

It is important to compare the outlined bilateral trading scheme with a basic price-based scheme. A cost-effective and environmentally safe price signal is a solution of the dual model to the basic primal model (5.3)–(5.4). It involves finding the price λ maximizing the following concave and, in general, non-differentiable function

$$\phi(\lambda) = \min_y \sum_{i=1}^n (f_i(y_i) + \lambda y_i).$$

A price signal λ decentralizes the solution of internal minimization problem into individual subproblems: find solutions $y_i(\lambda)$ minimizing functions $f_i(y_i) + \lambda y_i$. In general, solutions $y_i(\lambda)$ do not satisfy the balance equation, i.e., $\sum_{i=1}^n y_i(\lambda) \neq 0$, therefore the price λ has to be adjusted towards the desirable balance. The common idea is to change current λ_k at time $k = 0, 1, \dots$ proportionally to the imbalance, i.e., $\phi'(\lambda)$ for continuously differentiable $\phi(\lambda)$:

$$\lambda_{k+1} = \lambda_k + \rho_k \sum_{i=1}^n y_i(\lambda_k)$$

with a small step-size ρ_k . From the convergence results of quasi-gradient methods (see, e.g., discussion in [5]) it follows that with $\rho_k = \text{const}/k$, the sequence λ_k

converges to a price $\bar{\lambda}$ maximizing $\phi(\lambda)$. If $y_i(\lambda)$ are unique solutions (for any λ and $i = \overline{1, n}$), then $\phi(\lambda)$ is continuously differentiable function and $\bar{\lambda}$ fulfills the balance equation ($\sum_{i=1}^n y_i(\bar{\lambda}) = 0$) independently of the convexity $f_i(y_i)$. Otherwise, additional coordinated search is required to select \bar{y}_i from the set of solutions $y_i(\bar{\lambda})$ in order to guarantee (achieve) the balance $\sum_{i=1}^n y_i(\bar{\lambda}) = 0$. The coordination of parties is also required for tracking values of imbalances $\sum_{i=1}^n y_i(\lambda_k)$ for adjusting prices λ_k , $k = 0, 1, \dots$. Fundamental difficulties arise in the case of markets uncertainties (see Sect. 5.7) and duality gap, i.e., when $\max \phi(\lambda) < \min_y \{F(y), \sum_{i=1}^n y_i = 0\}$ for nonconvex function $F(y)$.

5.6 Computerized Multi-agent Decentralized Trading System

The proposed perfect market system implies that trades being bilateral, sequential (dynamic) and random do not impair the cost savings even if parties only have information on their own cost. However, there are essential obstacles that can inhibit real markets from perfect functioning according to proposed procedure. In a perfect market, a party that has sold permits in an early stage of the trading process would be able to cancel its earlier transaction. In the real emission trading market, this type of counter-actions may be impossible due to irreversibility of decisions: investments may already have been made, and these investment costs are largely sunk costs. This is the fundamental obstacle involved in the design of cost-minimizing and environmentally safe emission trading markets.

Price-based trading schemes have additional inherent obstacles. Designing environmentally safe and cost-effective price-based emission trading markets is equivalent to solving of the dual model asking for the same full information as the solution of the primary model. Additional critical limitation is the duality gap which occurs in nonconvex cases and uncertainty of market prices. The available computer technology and numerically stable optimization procedures allow to organize computerized (say, web-based) multi-agent decentralized trading system to resolve these issues.

One can imagine a distributed computer network that connects computers of parties with the computer of a central agency. The party in an anonymous manner stores information on its specific cost functions, and other characteristics of the underlying optimization model (5.8), (5.9) including specific probability distributions. The central agency stores information on the emission detection model. The computer of the central agency generates a pair of parties i, j and in an anonymous manner negotiates with computers of these partners a proper Δ_k that solves the subproblem (5.11). This can be easily organized without revealing private information of the parties. The process is repeated until equilibrium levels have been reached. This procedure allows to discover equilibrium solution that can then be implemented in reality. The information about the equilibrium price λ^* allows also to identify so-called core solution defining stable coalition of parties (see the Appendix). A network of interconnected computers is essential for a rapid, smooth and robust functioning

of emission trading market. There would also be a clear separation between a first stage, in which provisional bids are made between computers of parties and reconstructing is allowed, and a second stage, when contracts have been concluded and investments in emission control are implemented.

It is well known [2] that generally the market does not generate desirable outcomes if market prices fail to reflect socio-economic and environmental impacts. In this case it is typically necessary to establish negotiation processes between involved parties to determine desirable collective solutions. From this perspective, the proposed trading system can be viewed as a device for collective negotiations and decision-making in the presence of inherent uncertainties and irreversibilities.

5.7 Myopic Market Processes

The basic model (5.3), (5.4), (5.8), (5.9) takes long-term perspectives on emission permit trading. Parties use expectations and safety constraints in order to achieve cost-effective and environmentally safe outcomes robust against future developments. The resulting trading scheme is similar to non-monetary exchange economy [28] important for environmental control. There are no demand and supply functions. Instead, the safety constraints enforce parties to invest in emission and uncertainty reductions and consequently act as supplier of mutually beneficial emission permits until a global equilibrium emerges.

The situation becomes dramatically different in the case of price-based schemes under markets uncertainties affecting cost functions c_i , d_i of parties. The short term market perspectives orient parties on instantaneous information about prices and costs. At time interval k parties observe market-related components of uncertainty ω_k and thus know their instantaneous cost functions $c_i(x_i, \omega_k)$, $d_i(x_i, \omega_k)$. Based on this information, parties calculate cost functions

$$f_i(y_i, \omega_k) = \min_{u_i, x_i} [c_i(x_i, \omega_k) + d_i(u_i, \omega_k)], \quad (5.12)$$

subject to the safety constraints (5.8) conditional on observable uncertainties. They minimize then

$$\sum_{i=1}^n f_i(y_i, \omega_k) \quad (5.13)$$

subject to

$$\sum_{i=1}^n y_i = 0$$

by using observed price signals π_k , $k = 1, 2, \dots$, which separates joint model (5.13) into independent individual minimization of cost functions

$$f_i(y_i, \omega_k) + \pi_k y_i$$

by parties $i = 1, \dots, n$, where $\pi_k y_i$ is the cost of buying ($y_i > 0$) or selling ($y_i < 0$) emission permits y_i . If π_k coincides with an equilibrium spot price, then solutions $y_i(\omega_k)$, $i = 1, \dots, n$, of these individual models may coincide (Sect. 5.5) with the solution of joint model (5.13). In particular, they may satisfy the balance equation $\sum_{i=1}^n y_i = 0$. Otherwise, market prices π_k may cause disruptions of the balance and crashes of prices similar to the European carbon prices in April 2005. The following example illustrates a typical situation.

Example 5.3 (Market's uncertainty). Suppose that $f_i(y_i)$ are known deterministic functions, i.e., only prices are random. At time interval k there is a favorable situation for the exchange of emission permits for some parties i and j , e.g., $f'_i(y_i^k) \neq f'_j(y_j^k)$. Instead, high market price π_k , $\pi_k > \left| f'_j(y_j^k) \right| > \left| f'_i(y_i^k) \right|$, forces both parties to reduce emissions in the excess of targets in order to sell surpluses on the market. Disequilibrium price π_k creates an oversupply of emission permits that pushes the market price π_{k+1} towards 0. This may prevent to sell reduced emissions which turned to be of higher marginal costs with respect to new prices.

The myopic model (5.12)–(5.13) yields decisions $x_i(k, \omega_k)$, $u_i(k, \omega_k)$, $y_i(k, \omega_k)$, $k = 1, 2, \dots$. Such decisions depend on case-specific realizations of the random variable ω_k , therefore are not robust. At time interval $k + 1$ new observation ω_{k+1} may contradict ω_k requiring significant revisions of these decisions, which may be impossible due to their irreversibility. In order to achieve a convergence, the parties must adopt a precautionary incremental behavior with respect to arriving new information. Let us consider first this type of trading scheme for the basic model (5.3)–(5.4), (5.9).

The bilateral dynamic trading process of Sect. 5.5 has deep roots in the structure of so-called stochastic gradients in the linear subspace defined by (5.4). Namely, it is easy to prove that the vector

$$g(y) = (f'_1(y_1) - \frac{1}{n} \sum_{j=1}^n f'_j(y_j), \dots, f'_n(y_n) - \frac{1}{n} \sum_{j=1}^n f'_j(y_j))$$

is the projection of $gradF(y) = (f'_1(y_1), \dots, f'_n(y_n))$ in this subspace. A stochastic gradient then can be defined as the following. Pick up at random a pair (i, j) and define stochastic vector

$$\xi(y) = \frac{(n-1)}{2} (0, \dots, 0, f'_i(y_i) - f'_j(y_j), 0, \dots, 0, f'_j(y_j) - f'_i(y_i), 0, \dots, 0),$$

i.e., $\xi(y) = (0, \dots, 0)$ for $i = j$. This vector is a stochastic gradient of $F(y)$ [5], i.e., the conditional expectation $E[\xi(y)|y] = g(y)$, assuming that pairs (i, j) of distinct parties are chosen with equal probability $1/n(n-1)$. Therefore, instead of complete minimization of function (5.11) at step k , parties can move from y^k in the random direction $\xi(y^k)$ with a small step size α_k . This type of stochastic decentralized optimization processes are important in cases when functions $f_i(y_i)$

are not calculated exactly, e.g., they are affected by unknown random variables ω as in (5.12). It is easy to check that

$$F(y^k - \alpha_k \xi(y^k)) = F(y^k) - \alpha_k [f'_{i_k}(y_{i_k}^k) - f'_{j_k}(y_{j_k}^k)]^2 + o(\alpha_k), \quad (5.14)$$

where (i_k, j_k) is picked up at step k random pair (i, j) . Therefore, sequential change of emissions defined by equations

$$y^{k+1} = y^k - \alpha_k \xi(y^k) \quad (5.15)$$

produces monotonically decreasing for small α_k (contrary to standard stochastic gradient methods [5] random sequence $\{y^k\}$). The convergence analysis of this scheme is similar to the proof of the Theorem in the Appendix. It is also possible to derive from (5.14), that the scheme of Sect. 5.5 is in fact equivalent to the procedure (5.15) with the full step-size α_k equalizing marginal costs,

$$f'_i(y_i^{k+1}) = f'_j(y_j^{k+1}).$$

The basic adjustments in (5.15) are again pair-wise, but random encounters. Other encounters are also possible assuming that each party meets every other party.

This type procedure is also applicable in the case when parties use only observable random functions $c_i(x, \omega)$, $d_j(u, \omega)$. Suppose that instead of myopic decisions $y_i(k, \omega_k)$ parties make precautionary incremental and adaptive adjustments of vector y . We can define stochastic vector $\xi(y, \omega)$ similar to vector $\xi(y)$ as

$$\begin{aligned} \xi(y, \omega) = & \frac{(n-1)}{2} (0, \dots, 0, f'_i(y_i, \omega) - f'_j(y_j, \omega), \dots, f'_j(y_j, \omega) \\ & - f'_i(y_i, \omega), 0, \dots, 0) \end{aligned}$$

and proceed with changes of emission permits y^k according to procedure (5.15) with $\xi(y^k)$ substituted by $\xi(y^k, \omega_k)$. Under standard assumptions the conditional expectation $E[\xi(y, \omega)|y] = g(y)$, where $g(y)$ is the projection of $grad G(y)$,

$$G(y) = \sum_{i=1}^n E f_i(y, \omega). \quad (5.16)$$

The convergence of the trading scheme (5.15) with the vector $\xi(y^k, \omega_k)$ to a solution minimizing $G(y)$ subject to $\sum_{i=1}^n y_i = 0$ can be derived from general results on the convergence with probability 1 of stochastic quasigradient methods [5]. In particular, it requires the proper step-size multipliers α_k , e.g., $\alpha_k = const/k$ is applicable. This requirement presumes no knowledge of underlying data. Yet, it suffices to stabilize exchanges y^k .

Remark 5.5 (Short-term market decisions). Method (5.15) with vector $\xi(y^k, \omega_k)$ leads to an array of bilateral trading schemes with a variety of trading rules. Yet, it is

important to note a significant difference between functions $G(y)$ and $F(y)$ defined by (5.16) and (5.9), respectively; function $G(y)$ focuses on the variability of short-term market decisions $x_i(k, \omega)$, $u_i(k, \omega)$, whereas $F(y)$ focuses on forward-looking decisions x_i , u_i , y_i robust against all future eventualities ω . Thus, in contrast to trading scheme of Sect. 5.6, the minimization of function (5.16) takes short-term market perspectives on emission and uncertainty reductions driven by random observations ω_k of market situations. It also treats the safety constraints conditionally on myopic decisions $x_i(k, \omega)$, $u_i(k, \omega)$, that prevents to achieve robust cost-effective and environmentally safe outcomes specified by (5.3), (5.4), (5.7), (5.9). Such outcomes are guaranteed only through replacement of myopic overacting decisions $x_i(k, \omega)$, $u_i(k, \omega)$ by incremental adjustments of x_i , u_i similar to adjustments of y^k .

5.8 Concluding Remarks

The paper analyzes cost effective and environmentally safe carbon trading schemes explicitly incorporating different types of exogenous and endogenous uncertainties on emissions and the abatement costs. The feasibility of decentralized market pollution control mechanisms is usually discussed under strong assumptions that all actions are made simultaneously at known equilibrium prices, what implies existence of perfectly informed social planner. The examined dynamic bilateral trading schemes are not based on price signals, and the emerging emission prices implicitly depend on the costs functions and the safety constraints on environmental targets. With the safety constraints, the parties set the level of their exposure toward uncertainties and risks. The safety constraints discount the reported emissions to undershooting detectable levels. This type of undershooting concept or discounting should become a key element in a robust regulation of emission trades together with unified approaches to modeling emission uncertainties and cost functions. The paper shows that myopic price-based trading schemes are not able to achieve cost-effective and environmentally safe solutions. The irreversibility of trades calls for the use of the proposed computerized emission trading system providing, in a sense, collective decentralized regulation of trades. The procedures provide a constructive and easy approach for designing robust emission trading schemes. All decisions are fully decentralized, individual contrary to cost effective and environmentally safe price mechanisms requiring additional coordination to stabilize leading otherwise to nowhere trading processes. Our approach is close to important ideas on decentralized non-monetary exchange. Bilateral trading scheme with deterministic interval uncertainty has been applied [14] for the fossil fuel related carbon emissions of the major Parties of the Kyoto Protocol. Numerical findings indicate that the compliance costs increase significantly for USA, Japan and the European Union, if uncertainty of the emission levels is considered. However, although the Central and Eastern Europe, Russia, and Ukraine have larger uncertainties in emission levels, their net costs may decrease as they can sell emission reductions at a higher price. Additional simple calculations according to Remark 5.3 show that stochastic uncertainty in the emission levels reduces the compliance costs of parties.

Appendix

The convergence of trading scheme in Sect. 5.5 was proved in [8] for convex functions $f_i(y_i)$, $i = \overline{1, n}$. The following proof covers the case of nonconvex functions.

Proof of Convergence

Theorem (Convergence to an equilibrium). Let $f_i(y_i) \geq 0$ be continuously differentiable functions and let Y^* be the set of equilibriums equalizing marginal values f'_i as in (5.5), $F(Y^*) = F(y)$, $y \in Y^*$, y^k is defined as in Sect. 5.5. Then either

1. $y^k \in Y^*$ after a finite number of steps, or
2. The sequence $\{F(y^k)\}$ converges to its equilibrium value from the set $F(Y^*)$ and all cluster points of $\{y^k\}$ belong to Y^* , or
3. If Y^* contains only a single point y^* , then $\{y^k\}$ converges to this point.

Proof. The sequence $\{F(y^k)\}$, $k = 1, \dots$, is monotonically decreasing, $F(y^k) \geq 0$. Therefore, there exist a limit $\overline{F} = \lim_k F(y^k)$. Let us prove that $\overline{F} \in F(Y^*)$. Suppose there exists a convergent subsequence y^{k_s} , $y^{k_s} \rightarrow \overline{y}$, $s \rightarrow \infty$ and $\overline{y} \notin Y^*$. Therefore, there exist i, j such that $f'_i(\overline{y}_i) \neq f'_j(\overline{y}_j)$. It means that $\lim F(y^{k_s+1}) < F(\overline{y})$, what contradicts the convergence of $F(y^k)$, i.e., $\overline{F} \in F(Y^*)$, and all cluster points of the bounded sequence y^k belong to Y^* . Hence, if Y^* is a singleton, then y^k converges to y^* .

A Core Solution

From (5.10) it follows that at each step k cooperating parties i, j can redistribute joint cost $f_i(y_i^{k+1}) + f_j(y_j^{k+1}) = \phi_i^{k+1} + \phi_j^{k+1}$, $\phi_i^{k+1} < f_i(y_i^k)$, $\phi_j^{k+1} < f_j(y_j^k)$. Therefore at the equilibrium $y^* = (y_1^*, \dots, y_n^*)$ parties will deal actually with payments $\phi_i^* < f_i(y_i^0)$ such that $\sum_{i=1}^n \phi_i^* = \sum_{i=1}^n f_i(y_i^*) := F_I$ where $I = 1, \dots, n$. From this equation follows the Pareto efficiency of $\phi^* = (\phi_i^*)_{i=1, \dots, n}$. An important question is whether the grand coalition I of parties is stable, i.e., $\sum_{I \in C} \phi_i^* \leq F_C$ for any other coalition $C \subseteq I$. Accordingly, a distribution of payments ϕ^* is a core solution if it satisfies these two equations. The bilateral trading procedure allows to find the equilibrium price λ^* . If function $F(y)$ is convex, then the payment distribution $\phi_i^* = f_i(y_i^*) + \lambda^* y_i^*$ is a core solution. If the function $F(y)$ is globally Lipschitz continuous, then the core solution remains the same (see discussion in [11]).

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Chapter 6

Robust Design of Networks Under Risks

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Abstract Study of network risks allows to develop insights into the methods of building robust networks, which are also critical elements of infrastructures that are of a paramount importance for the modern society. In this paper we show how the modern quantitative modeling methodologies can be employed for analysis of network risks and for design of robust networks under uncertainty. The approach is illustrated by an important problem arising in the process of building the information infrastructure for the advanced mobile data services.

We show how the portfolio theory developed in the modern finance can be used for design of robust provision network. Next, the modeling frameworks of Bayesian nets and Markov fields are used for the study of several problems fundamental for the process of service adoption such as the sensitivity of networks, the direction of improvements, and the propagation of participants' attitudes on social networks.

6.1 Introduction

This paper is dedicated to a study of the network risks which are the key issues defining the robustness of infrastructures. There are similarities between network risks and catastrophic risks: both have interdependencies in space and time. An appropriate analysis of these two classes of risks requires adaptation, integration, extension and further development of methodologies for quantitative modeling of uncertainty and risks. Such methodologies have emerged during recent decades in diverse fields such as economics and finance, optimization, simulation of stochastic and multiagent systems. For a more detailed treatment we have selected two of

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such methodologies: portfolio theory of finance and Bayesian networks, both coupled with optimization approach show how these methodologies can be extended and applied for the study of network risks with the emphasis on information infrastructure.

Besides clarifying the methodological issues, we also aim at creation of integrated modeling decision support environment for analysis of network risks. Such environment enables identification and evaluation of critical bottlenecks inherent in important infrastructures seen as specialized networks and allow to give advice to planning and regulating bodies on robust design and improvement of these infrastructures.

More specifically, we look at the risk adapted performance networks composed of nodes and links of different levels of complexity. The risk adjusted performance of each node can be improved by selecting appropriate control parameters. In addition, performance of each node is affected by uncertainties. These network elements are designed from the point of view of local tradeoff between local performance and risk. The risk can be exogenous to the network as well as endogenous, generated by inappropriate functioning of other network elements. It is important that the overall performance of the network is also affected by the risk on the global level. This risk is understood as a possibility that the global performance can differ, sometimes drastically, from the expected network performance. The key issue in the designing of robust networks is to assure that a local level of risk/performance tradeoff results in a desirable risk/performance tradeoff on the global level. This is one of the central issues which we aim to clarify.

Here we briefly discuss three examples of infrastructure which can be described as risk/performance networks.

1. *Energy infrastructure.* Electric power infrastructure can be described as a network with several levels of hierarchy where the nodes correspond to production, distribution and consumption facilities while the arcs represent transmission lines. On the local level each node has production and consumption targets subject to uncertainty and risk manifested as the equipment failures, local demand variations, local weather patterns important for hydro and wind generation. On the global level this infrastructure should meet electric power demand of consumers and industries subject to uncertainties and risk of disruptions, prices for fuel and energy, weather, societal attitudes towards certain generation technologies and climate change.

Earlier such performance/risk tradeoff was much easier to achieve because the public utilities managed generation and transport in almost each country. Now the electric power industry is deregulated or being deregulated in almost all developed countries. The industry is now composed of many independent actors which decide their production plans according to the market conditions. Besides, the new types of actors have entered the field, like energy contract traders and speculators, and energy exchanges. Yet they have to act in concert if this infrastructure is to fulfill in a robust manner the energy needs of society at large. This is a critical issue as the power shortages in California and price surges in Norway have shown.

What are robust risk management methods which will mitigate these new risks which result from market forces and individual actors' behavior? What is the robust

way to assure that the local decisions on risk/performance tradeoff which every actor takes will transform into optimal or even acceptable tradeoff on the global level? What lessons developing countries can learn from the experience of developed countries in this respect? These are the questions which our paper aims to answer.

2. *Gas transport and distribution infrastructure.* Similar issues of networked risk/performance management arise in other types of infrastructure. For example, developing the gas transport and consumption infrastructure in Europe largely follows deregulation patterns of electric power infrastructure according to EU directives, and the same is true for the railroad transport.

3. *Information and communication network infrastructure.* It can be described as a superposition of several layers of hierarchical networks each one consisting of nodes connected with links. There are also mappings connecting different layers. The network nodes are represented by heterogeneous devices like routers, switches, cross-connects, etc. Each of these devices is equipped with control structures that govern communication flows through the network; the control is composed of communication protocols, routing tables, call admission rules, etc. The control parameters are tuned largely independently in order to meet performance targets of each node. Uncertainty on the node level comes from highly variable communication flows, but also from actions of adjacent nodes. There are also risks of equipment failures, congestion, malicious attacks, link failures which threaten the performance targets.

Each of the nodes is built to achieve admissible tradeoff between performance, costs and risk on the local level. The entire network, however, should satisfy various global performance targets, like a satisfaction of communication and information demand with quality of service guarantees for the user population. Besides, the operation of this infrastructure should be economically attractive for industrial actors which own its different parts. Global sources of risks and uncertainty include both external component (changing usage patterns and global malicious disturbances) and internal component (connected with conflicting interests of different actors).

How this local tradeoff between performance and risk at the node level affects the global tradeoff on the infrastructure level and vice versa? What are the economically sound principles for further robust development and operation of this infrastructure under inherent risk and uncertainty? Where are the bottlenecks which threaten its global performance? These are the questions which require the new methodology of the network risk analysis and robust risk management to which this paper aims to contribute.

Methods for taking optimal decisions under uncertainty and related issues of risk management have been at the center of methodological development in the couple of last decades, and recently they have met also a considerable and rising industrial interest. One should mention stochastic programming which is a hot topic in operations research community now and it has become an important modeling tool in quantitative finance, energy, telecommunications and other industrial fields. Understanding importance of risk management in finance resulted in the development of several risk management paradigms and industrial standards which are now being

gradually adopted also in other industrial branches. However, these and other methods mainly consider control of relatively simple systems facing external sources of risk and uncertainty. The real challenge is to look at the network of such systems and study the effects of risk and uncertainty on the network overall performance.

In this paper we look how such general considerations of network risks are adapted to the case of information infrastructure consisting of several components that are in different stages of development. We consider development and deployment of such a component characterized by increasing an importance: namely advanced mobile data services. We show how the related risks can be analyzed using and further extending for this case the quantitative risk modeling methodologies. In particular, the portfolio theory developed in finance is extended for analysis and design of service provision networks (Sects. 6.2–6.6). Next, Bayesian nets and Markov fields models are used in order to predict and analyze the sensitivity of networks, the directions of improvements, and the service adoption patterns, all of which depend on complex interplay of attitudes of different groups of population. Finally, we discuss development of two components of advanced methodological toolbox for analysis of network risk.

6.2 Cooperative Provision of Advanced Mobile Data Services

Design of advanced mobile data services to be carried on the 3G networks and the networks of next generations is the hot topic in telecommunication industry and research. This is because the business success of such services provision will define the business success of the mobile operators and other relevant industrial actors in the near to medium future. In this respect considerable attention is given to design and development of service provision platforms which support a set of tools and basic services that facilitate development, deployment and customization of specialized services by service providers and even non-professional end users.

Deployment and operation of service provision platforms and provision of individual services requires collaboration of different industrial actors contributing their individual capabilities and expertise to the common goal. One can think about fixed network operators, mobile operators, providers of different information content, internet providers, software developers and other actors that join forces to provide a successful service. Provision of a service involves assuming different roles and industrial actors can combine such roles. All this gives a rich picture of service provision environment where a multitude of actors cooperate and compete in order to deliver to customers a wide range of services in a profitable manner.

Understandably, the main research and development effort so far has been concentrated on technological and engineering aspects enabling provision of advanced mobile data services. The history of information technology testifies, however, that the possession of the best technological solution is not necessarily enough to assure the business success of an enterprise. A very important and sometimes neglected aspect is the design and evaluation of appropriate business model to support the

service provision. Business models for provision of a service requiring a single actor are pretty well understood, both organizationally and economically. This is the case, for example, of the provisioning of the traditional voice service over a fixed network. When an actor evaluates the economic feasibility of entering the provision of such service, he can employ quantitative tools developed by investment science, like estimation of the Net Present Value of such project [21]. Usually an actor should choose between several service provisioning projects, each providing return on investment and generating the risky cash flows. Then the portfolio theory [24] suggests the way to balance between return and risk and select the best portfolio of projects taking into account the actor's risk attitudes. The adequate risk management is especially important in a highly volatile telecommunication environment and the industrial standards in this respect are starting to emerge, originating from the financial industry [3]. Industrial projects in high-tech industries are often characterized by considerable uncertainty and at the same time carry different flexibilities. The real options approach [29] allows to take these flexibilities into account while making evaluation of the profitability of the project. Stochastic programming [2, 10, 13, 20] provide the optimization models for adequate treatment of uncertainty in the planning of service provision.

Business models for cooperative service provision involving different constellations of actors are studied to a much lesser extent. The understanding of their importance has lead to some qualitative analysis methods, e.g., [11, 19]; however, quantitative analysis at a level available for the single actor case remains a challenge. The methods mentioned above are developed to be used by a single actor engaged in selection and risk management of his portfolio of industrial projects. The influence of the other actors is present only implicitly on the stage of estimation of the future cash flows. This is not enough for an adequate analysis of collaborative service provision. Suppose, for example, that a service provider delivers a service to a population of users and receives a revenue for this delivery. If a service is composed of modules and enablers provided by different actors then this service provider has to decide on the revenue division between the actors which will make it attractive to them to participate in the service provision. The provider should explicitly incorporate in the evaluation of profitability of his project such revenue sharing decision together with a concept of what is attractive to other actors. Here we contribute to the adaptation and further development of the methods of evaluation and risk management of business models and industrial projects for the case of the collaborative service provision. We look at the actors engaging in a service provision as making a decision about the composition of their portfolio of services to which they are going to contribute. They do this independently following the risk management framework of the portfolio theory. The pricing and revenue sharing schemes induce each actor to contribute the right amount of provision capacity to its participation in the service provision. We develop a two tier modeling framework which supports selection of pricing and revenue sharing in an optimal way. This is done by utilizing the approach of stochastic optimization with bilevel structure [1] and combining it with the portfolio theory.

6.3 Simplified Model of the Service Portfolio

In this section we develop a quantitative description of the service provisioning model involving several actors having as the background the environment presented in the previous section.

6.3.1 Description of Services

A composition of a service can be quite complex, especially if we take into account that various components can be services themselves and subject to further disaggregation. For the purposes of clarity we start from a simple description which still possess the main features of the provision environment important for business modeling. Namely, two levels of the service composition will be considered here as shown in the example illustrated in Fig. 6.1.

In this case the service environment is composed of two types of services. The first type is comprised from services with structure and provision we consider in some detail. They can be provided in the context of a service platform and therefore they are referred to as *platform services*. There will be also *3rd party services* whose structure is of no concern to our modeling purposes. They are present in the model for the purpose of the adequate modeling of the environment in which the provisioning of the platform services happens. Let us now consider the model of provisioning of platform services.

The main building blocks of the platform services are service *enablers* indexed by $i = 1 : N$ and *services* indexed by $j = 1 : M$. Enablers are measured in units relevant for their description, like bandwidth, content volume, etc. The relation between enablers and services is described by coefficients λ_{ij} which measure the amount of enabler i necessary for provision of the unit amount of service j . Thus, a service j can be described by vector

$$\lambda_j = (\lambda_{1j}, \dots, \lambda_{Nj}) \tag{6.1}$$

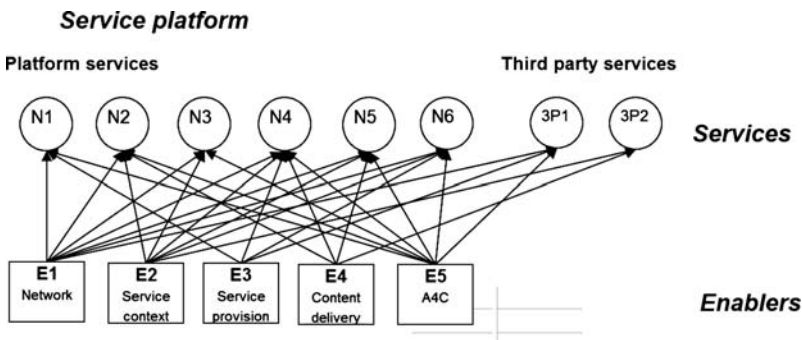


Fig. 6.1 Service provision for business persons on the move

This description is obtained from analysis of the usage scenarios described in the Sect. 6.2. A service j generates a revenue v_j per unit of service. This quantity depends on the service pricing which in its turn depends on the user behavior and market structure. For a moment let us assume that v_j is a random variable with a known distribution; in Sect. 6.4.3 we describe this revenue in more detail. The distribution can be derived from the expert estimates and from simulation models which would explore the structure of user preferences and market features. The random variables v_j can be correlated due to the service substitution, macroeconomic phenomena and other causes.

Services can be provided by different constellations of actors. In this paper we consider a constellation where the actors are the enterprises having the capability to provide service enablers assuming different *roles*, indexed by $k = 1 : K$. Actors may choose to join forces to provide a service. Contribution of a given actor consists of taking responsibility for provision of one or more enablers of the service. Sometimes these actors are referred to as *enabler providers*. There is also an actor providing the service aggregation functionality and organizing the overall service delivery to the end users; this actor is referred to as a *service provider*; it can provide the whole bundle of platform services and decides which services are included in this bundle. Often it collects the revenue from the end users and distributes it among the enabler providers.

Example 6.1. Service provision for business persons on the move (see Fig. 6.1). This is a simplified yet realistic example of service provision which was developed on the basis of the results of the EU project SPICE and the NFR project ISIS. The terminal service is a smart mobile phone used by a business person on the move.

We consider the services which run on the service platform and the third party services which partially compete with them, being accessible from the same terminal. We present a simplified example composed of just six native services, two third party services, and five enablers (in actual applications there are hundreds of services and dozens of enablers, distributed in several service platforms) available in this service platform. However, services in this platform correspond well to the business offer of a typical service provider. Service bundles have been defined in accordance with the market segments, corresponding customer classes, user behavior, requirements, and various subscription schemes. More specifically, we consider the following services.

Native services of the platform:

1. N1 – Messaging
2. N2 – Audio conferencing
3. N3 – Video conferencing
4. N4 – Location based services
5. N5 – News
6. N6 – Point of Interest service

Third party services:

1. 3P1 – Third party Information service
2. 3P2 – Third party News service

The following business actors collaborate in providing the mobile service bundle to the users:

1. E1 – *Network provider* – providing the network access.
2. E2 – *Context provider* – service context retrieval and management.
3. E3 – *Service provider* – responsible for service provision.
4. E4 – *Content provider* – content retrieval and management.
5. E5 – *Provider of A4C* (authentication, authorization, auditing, accounting and charging) enabler. This actor often coincides with the service provider, but one can envisage also the cases when it is a distinct actor.

Besides, there are one or more providers of the third party services which are in partial competition with the platform services.

The objective of an enabler provider is to select a portfolio of services to which this actor will make a contribution. This decision is made on the grounds of balance between projected profit from enabler provision balanced against the risk of variations in demand and service acceptance among the prospective users of services. In order to quantify such decision process it is necessary to use a simplified profit model for an actor.

It is assumed that the revenue v_j generated by a unit of service j is distributed among the actors participating in the service creation. There can be different schemes for such subdivision. It is assumed here that this distribution is performed using a vector of revenue shares

$$\gamma_j = (\gamma_{1j}, \dots, \gamma_{Nj}), \quad \gamma = (\gamma_{11}, \dots, \gamma_{N1}, \dots, \gamma_{1M}, \dots, \gamma_{NM})$$

such that an actor which contributes with the enabler i receives the revenue $\gamma_{ij}v_j$. Determination of these revenue sharing coefficients is one of the objectives of the design of the business model for service provision.

Besides platform services the actors can supply enablers also to the third party services. The structure of these services is not specified and it is assumed that they are fully described by the revenue v_{ij} generated by provision of the unit of enabler i to third party service j , $j = M + 1, \dots, \bar{M}$.

This example will be treated in more detail in Sect. 6.6.

6.3.2 Profit Model of an Actor

Let us consider the situation when all the actors have already developed the capacities for provision of enablers. Thus, the investment process necessary for creation and expansion of these capacities is not discussed here; however, it is considered at the later stages. For this reason at this stage it is enough to consider only the variable costs due to the operation of capacities and provision of enablers. Alternatively,

one can assume that the cost structure includes both the operational and discounted portion of the investment costs for enabler development, recalculated down to the enabler and the service instances.

For further formulation of the actor's profit model let us introduce the following notations:

c_{ik} – Unit provision costs for enabler i by actor k

W_{ik} – Provision capability of enabler i of actor k

x_{ijk} – The portion of provision capability for enabler i of actor k dedicated to participation in provision of service j

Now the revenue of actor k obtained from contribution to provision of the platform service j can be expressed as follows. The quantity $x_{ijk}W_{ik}$ is the volume of provision of enabler i dedicated by actor k to service j . Assuming that the required quantity of other enablers is available, this results in the volume of service j in which the actor k participates to be $x_{ijk}W_{ik}/\lambda_{ij}$. The total revenue from this service is equal to $v_j x_{ijk}W_{ik}/\lambda_{ij}$ and the part of the revenue which goes to actor k is $v_j x_{ijk}W_{ik}\gamma_{ij}/\lambda_{ij}$. For the third party service the revenue will be $v_{ij}x_{ijk}W_{ik}$.

The total costs incurred by actor k for the provision of enabler i to service j is equal to $x_{ijk}c_{ik}W_{ik}$.

In order to simplify the following discussion let us assume now that the actor k participates in the provision of service j by contributing only one enabler $i = i(k, j)$ or assuming only one role. Taking the profit π_k to be the difference between the revenue and costs, the profit of the actor k can be expressed as follows:

$$\begin{aligned}\pi_k &= \sum_{j=1}^M \left(v_j x_{ijk} W_{ik} \frac{\gamma_{ij}}{\lambda_{ij}} - x_{ijk} c_{ik} W_{ik} \right) + \sum_{j=M+1}^{\bar{M}} \left(v_{ij} x_{ijk} W_{ik} - x_{ijk} c_{ik} W_{ik} \right) \\ &= \sum_{j=1}^M x_{ijk} W_{ik} c_{ik} \left(\frac{v_j \gamma_{ij}}{c_{ik} \lambda_{ij}} - 1 \right) + \sum_{j=M+1}^{\bar{M}} x_{ijk} W_{ik} c_{ik} \left(\frac{v_{ij}}{c_{ik}} - 1 \right)\end{aligned}$$

In the expression above index i depends on the values of indices j and k . Now let us assume that the actor k assumes only one role which consists in the provision of enabler i to different services which require this enabler. Thus, we consider a generic actor whose role is to provide enabler i to different services. Then we can simplify notations by taking $x_{ijk} = x_{ij}$, $W_{ik} = W_i$, $c_{ik} = c_i$, $\pi_k = \pi_i$. In this case the profit is defined by:

$$\pi_i = W_i c_i \left(\sum_{j=1}^M x_{ij} \left(\frac{v_j \gamma_{ij}}{c_i \lambda_{ij}} - 1 \right) + \sum_{j=M+1}^{\bar{M}} x_{ij} \left(\frac{v_{ij}}{c_i} - 1 \right) \right).$$

Dividing the profit by the total costs $W_i c_i$ we obtain the return r_i on investment by a generic actor which assumes the role of provision of enabler i to services which

require this enabler:

$$r_i = \sum_{j=1}^M x_{ij} \left(\frac{v_j \gamma_{ij}}{c_i \lambda_{ij}} - 1 \right) + \sum_{j=M+1}^{\bar{M}} x_{ij} \left(\frac{v_{ij}}{c_i} - 1 \right). \quad (6.2)$$

6.3.3 Service Portfolio: Financial Perspective

The profit representation (6.2) allows us to look at the enabler provision from the point of view of the financial portfolio theory [24]. The actor with the role to provide the enabler i has to choose from all possible available services requiring this enabler the set of services which provide this enabler. In other words, he has to select his service portfolio. This portfolio is defined by shares x_{ij} of his provision capability,

$$x_i = (x_{i1}, \dots, x_{i\bar{M}})$$

Return coefficients associated with the participation in each platform service are expressed as

$$r_{ij} = \frac{v_j \gamma_{ij}}{c_i \lambda_{ij}} - 1, \quad j = 1 : M \quad (6.3)$$

and for the third party services these coefficients are

$$r_{ij} = \frac{v_{ij}}{c_i} - 1, \quad j = M + 1 : \bar{M}. \quad (6.4)$$

These coefficients depend on the random variables, in particular the revenue per unit of service v_j and the revenue per component provision v_{ij} . Randomness here is due to the uncertainty in demand and the user acceptance of service. Also the enabler provision costs c_i and enabler shares λ_{ij} are random variables due to the uncertainty inherent in the service usage patterns and the evolution of costs. Besides, the costs c_i are often estimates of the provision costs of enabler provider i made by another actor. Such estimates are inherently imprecise and therefore described by random variables similarly to how it was done in [1]. The expected return coefficients are

$$\mu_{ij} = \gamma_{ij} \mathbb{E} \frac{v_j}{c_i \lambda_{ij}} - 1, \quad j = 1 : M, \quad \mu_{ij} = \mathbb{E} \frac{v_{ij}}{c_i} - 1, \quad j = M + 1 : \bar{M} \quad (6.5)$$

and expected return $\bar{r}_i(x_i)$ of the service portfolio is

$$\bar{r}_i(x_i) = \sum_{j=1}^{\bar{M}} \mu_{ij} x_{ij} = \sum_{j=1}^M x_{ij} \left(\gamma_{ij} \mathbb{E} \frac{v_j}{c_i \lambda_{ij}} - 1 \right) + \sum_{j=M+1}^{\bar{M}} x_{ij} \left(\mathbb{E} \frac{v_{ij}}{c_i} - 1 \right) \quad (6.6)$$

However, the realized return can differ substantially from the expected return due to the uncertainty discussed above. This introduces the risk $R(x_i)$ for an actor which

assumes the enabler provision role. The financial theory traditionally measures this risk as the variance of the portfolio return [24]. Recently several different risk measures were introduced, in particular Value at Risk (VaR) and its many modifications. The VaR has attained the level of industrial standard in the financial risk management [3]. In this section the variance and the standard deviation of the return are used as the risk measure. It is not clear yet what is the best risk measure to consider in the context of portfolio problems related to the collaborative service provision. Besides the classical choice of standard deviation which we follow here, the promising candidates are the *VaR* mentioned above [14] and the Conditional Value at Risk [15]. The identification of the most relevant risk measure in this context is outside the scope of this paper; it will be pursued in our subsequent research. What is important here is the consideration of the risk measures allow an actor to estimate the probability and size of his future losses. Thus, we take

$$R(x_i) = \text{StDev}(r_i(x_i)) = \text{StDev} \left(\sum_{j=1}^{\bar{M}} r_{ij} x_{ij} \right) \quad (6.7)$$

where the return coefficients r_{ij} are taken from (6.3), (6.4).

Portfolio theory looks at the portfolio selection as the trade-off between risk and return. Its application to our problem of service portfolio consists of the following steps.

1. *Construction of the efficient frontier.* An average return target η is given. The risk of service portfolio is minimized with constraint on this return target. The risk minimization problem is defined by:

$$\min_x \text{StDev}^2 \left(\sum_{j=1}^{\bar{M}} r_{ij} x_{ij} \right) \quad (6.8)$$

$$\sum_{j=1}^{\bar{M}} \mu_{ij} x_{ij} = \eta \quad (6.9)$$

$$\sum_{j=1}^{\bar{M}} x_{ij} = 1, \quad x_{ij} \geq 0 \quad (6.10)$$

Solution of this problem for all admissible values of target return η provides the set of service portfolios which are a reasonable candidates for selection by the actor provides the enabler j . They constitute the *efficient frontier* of the set of all possible service portfolios. This concept is illustrated in Fig. 6.2.

Each service portfolio x can be characterized by the pair (risk, return) defined by (6.7) and (6.6) respectively. Therefore it can be represented as a point in the risk-return space depicted in Fig. 6.2. The set of such points for all possible portfolios describes all existing relations between risk and return and is called *the feasible set*. Which of possible service portfolios an actor should choose? It depends on the

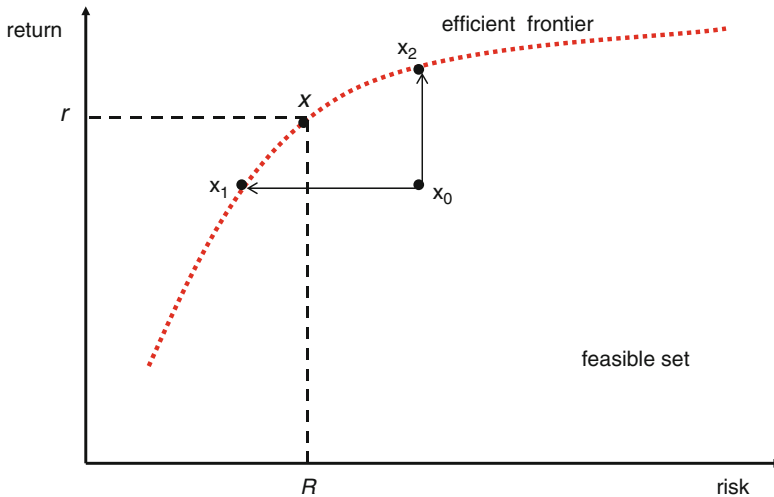


Fig. 6.2 Selection of service portfolio

objectives which an actor pursues. Here we assume that an actor's decision depends on return and risk only. Namely, an actor seeks the highest possible return among equally risky alternatives and the lowest possible risk among equally profitable alternatives. This is a simplification, because in reality the actors can be driven by other considerations, like increase of market share, revenue, regulatory constraints, etc. However, the consideration of only risk and return provides with a reasonable starting point for analysis of business models. More complex cases can be taken into account in a similar manner by introducing additional constraints on the feasible set or by modifying the concept of performance. For example, suppose that an actor has three objectives: the return and the market share to maximize and the risk to minimize. Then these three criteria will be used for specification of an achievement function used in the multi-criteria problem analysis, see, e.g., [30]. Such an analysis supports examination of efficient solutions having different trade-offs between the criteria values, and help the users in finding such trade-offs the correspond best to his/her preferences.

The example illustrated in Fig. 6.2 show that some service portfolios should be preferred to others. For example, let us consider portfolio x_0 to which corresponds the point in risk-return space inside the feasible set shown in Fig. 6.2. It is clear that the portfolio x_2 should be preferred to x_0 , because x_2 has the same risk as portfolio x_0 but it has larger return. Similarly, portfolio x_1 also should be preferred to x_0 because it provides the same return with smaller risk. In other words, portfolio x_0 is dominated by both portfolios x_1 and x_2 and therefore it is not rational to select it when only these two criteria are considered. The actor whose decisions are guided by risk and return should consider only nondominated portfolios which constitute the *efficient frontier*, depicted by dotted curve in Fig. 6.2. Such an efficient frontier can be computed by solving the problem (6.8)–(6.10) for different values of η . It can

be considered as a set of efficient solutions of the two objective optimization problem: minimizing risk and maximizing return under constraints (6.10). The choice of a solution from the efficient set depends on the preferences the user has between risk and return, and it is easy for experienced managers.

The above outlined approach to the multicriteria analysis is suitable for problems with two criteria, i.e., for which it is possible to compute and analyze the *efficient frontier*. For more than two criteria one should use a suitable method for multi-criteria analysis. However, one has to note that the most popular approach to multi-criteria analysis believed to be reliable and intuitive, namely the aggregation of criteria through a weighted sum is actually neither reliable nor intuitive. Limitations of the weighted-sum approach are discussed, e.g., in [22, 23]. Therefore, for analysis of problems with more than two criteria one should consider a truly multicriteria analysis approach, see, e.g., [30], and a modular tool (such as ISAAP described in [18]) that supports interactive analysis of trade-offs between conflicting objectives.

2. *Selection of the target service portfolio.* The previous step resulted in the selection of a much smaller set of efficient service portfolios from the set of all possible service portfolios. These portfolios form the efficient frontier in the risk-return space. An actor selects its target service portfolio from this efficient set by choosing the trade-off between risk and return. One way to achieve this trade-off is to consider the largest risk an actor is willing to take. Suppose that the value of such risk is R (see Fig. 6.2). Then the actor should choose the portfolio x on efficient frontier with this value of risk. Suppose that this service portfolio yields return η . No other portfolio yields better return without increasing the risk. If an actor is not satisfied with return η this means that he should increase his risk tolerance or look for opportunities to participate in the service provision not yet described in this model. Or, such actor should seek more advantageous revenue sharing scheme.

From these considerations it is clear that all important opportunities of participation in service provision should be included in the model. For example, suppose that an actor assumes the role of the content provision and can contribute a content to the advanced mobile data service, and at the same time the same content can be contributed to, say, traditional newspapers. Both opportunities should be included in the model with the traditional service being modeled as a third party service.

6.4 Modeling of Collaborative Service Provision

In the previous section we highlighted the importance of having an adequate forecast of the cash flows generated by services in order to quantitatively evaluate the economic future of the service, and business models which support the service provision. Due to the inherent uncertainty of the user response and technological development any such forecast should be given in terms of random variables which assign probabilities to different scenarios of user response and possible evolution of other uncertain parameters. Forecasts should take into account the mutual influence

of services which result in correlation between cash flows generated by different services.

Such a model allows to consider providers of different service enablers as actors which independently select the service portfolios having their targets described in terms of return on investment and risk tolerance. However, a service can become a reality only if the participation in its provision is consistent with these individual targets. This means that all actors which cover the roles indispensable for provision of a particular service should have the corresponding services in their efficient service portfolio. In other words, the service portfolios of the relevant actors should be *compatible*. There are several items which affect the risk/return characteristics of a service portfolio and decide whether a particular service is offered. One is the cash flow generated by a service j , another is the revenue sharing scheme γ_j . Besides, the enabler provision capacities, industrial risk/return standards, market prices, all play a role in making service portfolios compatible. In this section we characterize the properties which facilitate the service portfolio compatibility and develop a model for selection of the revenue sharing scheme.

6.4.1 Service Provision Capacities

According to (6.1) a platform service j is described by vector λ_j of the service enablers. Let us denote by I_j the set of enablers which are present in the service description in nonzero quantities:

$$I_j = \{i : \lambda_{ij} > 0\}$$

For each enabler $i \in I_j$ an actor willing to take the role of provision of this enabler should be found. This means that the position j in the service portfolio of an actor that provides enabler $i \in I_j$ should be positive: $x_{ij} > 0$. The value of this position allows to estimate the enabler provision capacity which an actor should possess. Indeed, x_{ij} is a fraction of provision capacity which an actor is going to dedicate to provision of enabler i to service j . Therefore, λ_{ij}/x_{ij} is the capacity necessary to provide a unit of service j . Suppose that B_j^{\min} is the minimal volume of provision of service j which makes such provision viable, and B_j is the target volume of service provision for a generic constellation of actors which is going to provide this service. Then we have the following constraints on the service provision capacities of actors:

$$W_i x_{ij} \geq \lambda_{ij} B_j^{\min}, \quad i \in I_j \quad (6.11)$$

if the provision of service j will be viable at all, and

$$W_i x_{ij} \geq \lambda_{ij} B_j, \quad i \in I_j \quad (6.12)$$

if only one actor with provision capability of enabler i is desirable in the constellation which provides service j . These constraints can help to make decisions about the nature of the actors which should be encouraged to participate in the provision of different services. For example, some enablers of some services will be provided by established actors with large provision capacity. In such cases the share x_{ij} of capacity dedicated to service j can be small. In other cases the service enablers will be provided by startups with relatively small capacity. In such cases the share x_{ij} should be smaller or equal to 1. These shares implicitly depend on the revenue sharing scheme γ_j through the solution of problem (6.8)–(6.10) and in the latter case it may be beneficial on the initial stages of service penetration to encourage startups by appropriate adjustment of the revenue sharing scheme.

The constraints (6.11)–(6.12) can be also looked at as the constraints on the composition of service portfolio. Suppose that W_i^{\max} is the maximal desirable component provision capacity which an actor providing enabler i should possess. Then the smallest share x_{ij} dedicated to service j should be

$$x_{ij} \geq \lambda_{ij} \frac{B_j^{\min}}{W_i^{\max}}, \quad i \in I_j \quad (6.13)$$

6.4.2 Risk/Return Industrial Expectations

Provision of advanced mobile data services involves different actors coming from different backgrounds and industries. There are many startups, but there are also established actors from other industrial branches. For example, the content provision where the same content can be provided to newspapers, internet and mobile terminals. Such actors have preferences for tradeoffs between admissible and/or desirable returns on investment and the corresponding risks. Often such preferences are influenced by industrial standards and expectations inherited from their previous activities. One way to express such expectations is to include all generic projects, in which an actor can be involved in its traditional business, as services in the set of all third party services considered in the model. This is especially useful approach, if the revenues from the traditional activities influence and are influenced by the revenues from the mobile services under consideration. Another possibility is to account for these expectations explicitly. This can be done by introducing the relation between the expected return $\bar{r}_i(x_i)$ and risk [26] $R(x_i)$ from (6.6), (6.7) as follows:

$$\bar{r}_i(x_i) \geq a_i + b_i R(x_i) \quad (6.14)$$

where a_i is the return on investment associated with traditional activity while $b_i R(x_i)$ is the risk premium associated with the participation in provision of the advanced mobile data services. The coefficients a_i and b_i will depend also on individual characteristics of an actor, e.g., its size, market position.

Additionally an actor has the risk tolerance expressed in terms of the upper bound on risk which it is willing to take irrespective of return:

$$R(x_i) \leq \bar{R} \quad (6.15)$$

The upper bound on admissible risk \bar{R} also depends on the characteristics of a particular actor. To put it simply, this is the maximal loss an actor can afford during the time period under consideration.

6.4.3 Pricing

The revenue per unit of service v_j together with the service composition λ_j and the revenue sharing scheme γ_j defines the unit price p_i of enabler i :

$$p_i = \frac{v_j \gamma_{ij}}{\lambda_{ij}}$$

This is a random variable since the revenue is random. Therefore the expected price $\bar{p}_i = \mathbb{E}p_i$ is:

$$\bar{p}_i = \frac{\gamma_{ij}}{\lambda_{ij}} \mathbb{E}v_j$$

An actor providing the enabler i may have the target p_i^* for the price of its product, and the tolerances Δ^+ and Δ^- within which it is willing to accept a price. These targets can result from the market prices in established industries, internal market studies, or internal cost estimates. This leads to the following constraint

$$p_i^* - \Delta^- \leq \frac{\gamma_{ij}}{\lambda_{ij}} \mathbb{E}v_j \leq p_i^* + \Delta^+ \quad (6.16)$$

This constraint should be taken into account while considering the revenue sharing schemes.

6.4.4 Revenue Sharing Schemes

Now let us look at the problem of selecting the revenue sharing coefficients γ_j which would be compatible with the concerted provision of a platform service. Summarizing the discussion presented in Sects. 6.3, 6.4.1 and 6.4.2 we obtain that the actor supplying enabler i selects a portfolio of services $x_i = (x_{i1}, \dots, x_{iM})$ by solving the following problem

$$\max_{x_i} \bar{r}_i(x_i, \gamma_j) \quad (6.17)$$

subject to constraints

$$\sum_{j=1}^{\bar{M}} x_{ij} = 1, \quad x_{ij} \geq 0 \quad (6.18)$$

$$\bar{r}_i(x_i, \gamma_j) \geq a_i + b_i R(x_i, \gamma_j) \quad (6.19)$$

$$R(x_i, \gamma_j) \leq \bar{R} \quad (6.20)$$

where $\bar{r}_i(x_i, \gamma_j)$ is the expected return of the actor on his expenditure and $R(x_i, \gamma_j)$ is the risk defined by (6.6) and (6.7), respectively. We emphasize here the dependence of risk and return on the revenue sharing scheme γ_j . Solution of this problem provides service portfolios $x_i(\gamma_j)$ for all generic actors providing enabler i for the platform services $j = 1 : M$. These service portfolios depend on the revenue sharing schemes γ_j . Let us now concentrate on a j -th service. In order that a provision of this service becomes possible it is necessary that all actors providing the necessary enablers to this service include it in their service portfolios in the desirable proportions. This is denoted by:

$$x_i(\gamma_j) \in X_j \text{ for all } i \in I_j \quad (6.21)$$

where the set X_j is defined, for example, by constraints (6.13). The feasible set of the revenue sharing coefficients is defined by the set X_j and if these conditions are not satisfied then the service is not available.

Suppose now that the enabler number 1 of service j is a service aggregation enabler which is provided by an actor which bears overall responsibility for the functioning of service and receives the revenue stream from the end users. Its responsibility includes also the division of the revenue stream between the participating actors and thus also a selection of the revenue sharing coefficients γ_j . It selects these coefficients in such a way that the constraints (6.21) are satisfied. Between all such revenue sharing coefficients selects those which maximize its return. This can be formulated as the following optimization problem.

$$\max_{\gamma_j} \bar{r}_1(x_1(\gamma_j), \gamma_j) \quad (6.22)$$

subject to constraints

$$x_i(\gamma_j) \in X_j \text{ for all } i \in I_j \quad (6.23)$$

$$\gamma_j \in \Gamma_j \quad (6.24)$$

where the set Γ_j is defined, for example, by constraints (6.16). Even simpler, this actor may wish to maximize its revenue share

$$\max_{\gamma_j} \gamma_{1j} \quad (6.25)$$

under constraints (6.23)–(6.24). Observe that the feasible set of this optimization problem depends on the solution of the other actor's optimization problems (6.17)–(6.20).

6.5 Properties of the Models and Implementation Issues

In Sect. 6.4.4 we presented two models for strategic assessment of collaborative provision of mobile data services. These models possess quite complicated structure, although we made a few simplifying assumptions. They can be considered as a special type of stochastic optimization problems with bilevel structure [1], where the lower level is composed of the problems of individual component providers (6.17)–(6.20) while the upper level contains the problem of service provider (6.22)–(6.24). Stochasticity comes from the uncertainty inherent in the characteristics of advanced data services and the user response to them. So far we have adopted a simple treatment of uncertainty by substituting the random variables by their expected values, and, when the special structure of the problem allowed, the covariance matrix. This can be viewed as a special type of the deterministic equivalent of the stochastic programming problems, a technique widely used in stochastic programming (see [2] for more discussion on different types of the deterministic equivalents). A more detailed description of the uncertainty can be introduced in these models following the approach presented in [1]. Moreover, different bilevel optimization problems have drawn considerable attention recently, see [5, 6, 25, 28].

Such problems provide quite a challenge to the state-of-the-art numerical optimization procedures. While many theoretical issues are understood reasonably well, the solution techniques have not yet reached the off-the shelf commercial applicators available for linear and some nonlinear programming problems. The main challenge here is that the upper level problems can be highly nonlinear and nonconvex with multiple local minima. Therefore a substantial work is needed for exploiting the structure of such problems. Still, our aim here is to describe a set of decision support tools for evaluation of business models, where the computational complexities should be hidden from the end user. We have found that this aim can be achieved by a combination of customized implementation with the use of general purpose mathematical modeling systems and commercial software. The general architecture of the system under development is shown in Fig. 6.3.

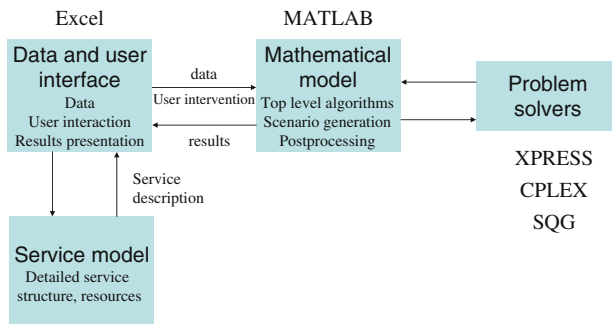


Fig. 6.3 Architecture of decision support system for evaluation of business models of service provision

The system is composed of four components: data and user interface, a set of service models, a set of mathematical models, and a library of solvers.

Data and user interface is implemented in Excel due to its familiarity to potential users. Its purpose is to provide an easy tool for storing and changing the data describing the service and customer properties, for presentation of results of business modeling and for providing the capability to the system user to ask what-if questions pertaining to different scenarios. For example, the efficient frontier illustrated in Fig. 6.2 is presented to the user through this component.

Service model provides the capability to perform detailed modeling of advanced data services. It is implemented in a specialized modeling language which has the features necessary for describing communication sequences. This model provides the aggregated description of services composition λ_j from (6.1).

Mathematical model implements the quantitative description of the business decision process of collaborative service provision from the previous sections. It imports data from data interface and implements the top level structures and algorithms necessary for representation and solution of models (6.17)–(6.20), (6.22)–(6.24). The custom algorithms for analysis and solution of these models are also implemented in MATLAB. This component is also responsible for calling external software for solving subproblems where standard approaches and commercial software are available. For example (6.8)–(6.10) is a quadratic programming problem which can be solved by many solvers, e.g., by the CPLEX and MATLAB optimization toolboxes.

Library of solvers contains solvers for linear and nonlinear programming problems and some a specialized solvers for stochastic programming problems, e.g., SQG [12].

The system depicted in Fig. 6.3 is now in advanced stages of development, in particular the service model component and some mathematical models of service provisioning were implemented in MATLAB [1]. The next section describes some of the results of a case study performed using this system.

6.6 Case Study

This case study deals with analysis of the service provider centric business model for provision of the platform bundle of services to a business person on the move who uses a smart mobile phone to access the corresponding service offer. The setting of this case study is described in Example 6.1 introduced in Sect. 6.3. The prototype of the decision support system implementing models presented in this paper was used for the analysis of this case study; it includes the models described in Sects. 6.3 and 6.4.

Considerable data preparation effort was made for this case study. First of all, we have developed the service composition matrix, showing which enabler participates in which services. This relation between different enablers and services is shown in Fig. 6.1. We have obtained also an average estimate of the service usage (in service

instances) in a period of interest, and prices per service instance. We have estimated these data by averaging various service composition and business scenarios. On the basis of technical and economic analysis we have obtained the cost estimates and the correlation matrix showing the correlation between the usage of services and the variance of service usage.

Suppose that the service provider uses our DSS for performing the feasibility study for provision of this bundle of business services similarly to the discussion at the end of Sect. 6.3.3. There are many different what-if questions of interest to the prospective service provider to which this DSS can provide answers. Let us provide an example of this analysis. Suppose that a manager of the service provider feels that the success of the whole enterprise depends critically on the quality and offer of specialized content which can be obtained for his services by engaging prospective content providers (enabler E4 from Fig. 6.1). He wants to get an insight into the properties of the content providers which may be interested in collaboration with the provider and in the chances that his service offer in this respect will stand against the competition of the third party services. One way to do this is to look how the service portfolios and risk/profit preferences of prospective partners depend on correlation and relative pricing of his offer against the offer of competition. Figures 6.4 and 6.5 provide examples of answers which our DSS can deliver.

Figures 6.4 and 6.5 show how the characteristics and attitudes of the content providers towards the service platform depend on the alternatives the competitors can offer to them. Figure 6.4 shows risk/profit efficient frontiers similar to the frontier presented in Fig. 6.2 while Fig. 6.5 depicts the percentage share of the content provision capability of the content providers dedicated to the service platform. In other words, the Fig. 6.5 shows the market share of the service platform in the market for this specific type of content provision dependent on the risk tolerance of the content providers. The competing offer is described by the average price per unit of

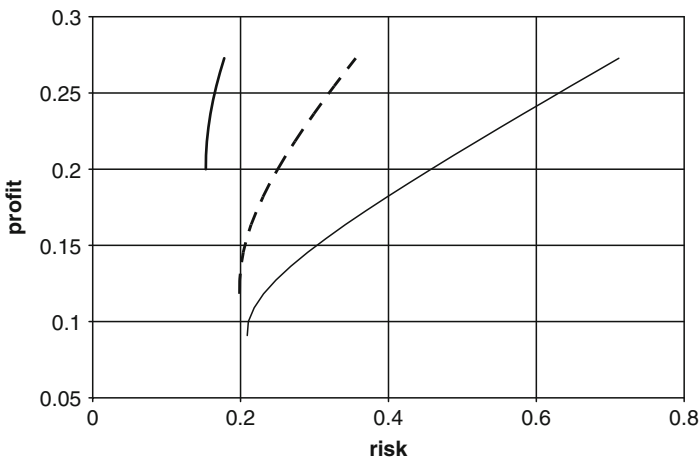


Fig. 6.4 Dependence of risk/return preferences of content providers on the strength of competition

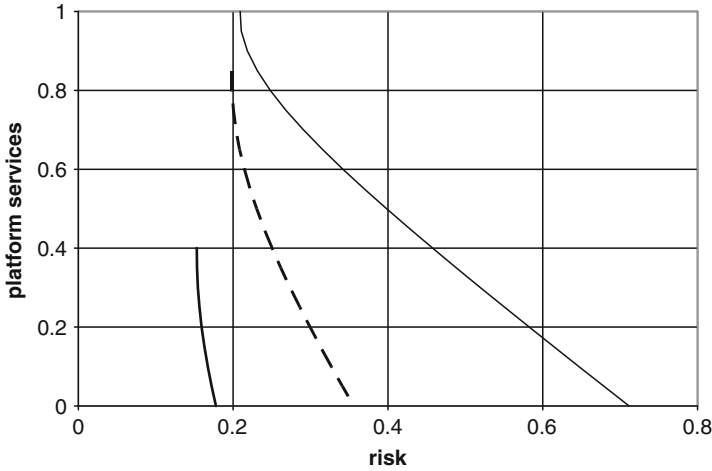


Fig. 6.5 Share of platform services in the service portfolios of content providers

content and by how the actual price can differ from the average price dependent on the future market conditions, as measured by the price variance.

The figures present three scenarios. In all scenarios the competition tries to undercut the service platform by offering about 15% higher average price to content providers for their services. The three scenarios differ by how strong the competition is, namely its capability to maintain the price consistently higher under the changing market conditions. In scenario 1 shown by the thick solid line the competition is strong and has its price variance about two times smaller compared with the platform offer. In scenario 2 depicted by the thick dashed line the competition is about as strong as the platform offer and has the similar price variance. In scenario 3 shown with thin solid line the competition is weaker than the platform and has about twice higher variability of its offer to content providers than the platform.

The results show that in scenario 1 with the strong competition only economically weak content providers with small tolerance towards losses are interested in the collaboration with the platform. Often this corresponds to small firms or even individuals who can not sustain large losses. For such entities participation in the platform means additional security and insurance against losses in the case the strong competing offer prove to be deceitful. Even then, the interest of such firms drops sharply when their risk tolerances grow even by a small amount.

Scenario 3 corresponds to the opposite case when the platform faces aggressive but economically relatively weak competition. Its weakness manifests itself in a large variability of the price offer to the content providers despite the 15% higher average price. In this case the market share of the platform services is much higher and the platform manages to attract also strong actors with higher capacities to sustain losses. Also the market share drops slower with the increase of the loss tolerance of the agents. Scenario 2 corresponds to the intermediate case when the competition

is about as strong as the platform and has about the same capability to maintain its price offer to the service providers.

Similar patterns arise when the variability of the revenue stream of content providers is not due to the changes in the unit price of the content but due to the variability of the usage frequencies of this content. Having these predictions, the platform service provider can now realistically weight its strengths and weaknesses, invest more effort into the market research and decide under which market conditions, with what kind of partners, and with what kind of competition it can successfully operate the platform.

6.7 Dynamics of Attitudes

In the previous sections we have analyzed the network risks which arise in the process of collaborative provision of advanced mobile data services which form an important part of information infrastructure. In this section we discuss another aspect of the same situation, namely the process of service adoption by a heterogeneous population of users. The process of service adoption is of fundamental importance to the successful development of information infrastructure because by its very nature the value of the infrastructure for a given user grows with the amount of users already covered by its different components. Adopting the language of microeconomics one can say that the elements of information infrastructure exhibit strong externalities. Modeling these externalities and related risks requires tools and approaches for quantitative modeling of attitudes. In this and subsequent sections we consider a possible methodology based on Bayesian nets.

We formulate a stochastic, dynamic model of attitude formation that takes special account of individual interactions and networks governing intrinsic dynamics of attitudes. The model also accounts explicitly for various external factors such as new information, stimulus, events, actions or some sort of social pressure. If different sets of external factors are activated at different times, the system may show more or less complex dynamics, in particular, it may lead to different alternative attractors. We distinguish two types of influences: (1) interactions among individuals, and (2) the influence of external factors.

According to our model different individuals may receive different information. Information with subjective judgments is transformed through chains of communications to other individuals. Attitudes change in a probabilistic manner depending on attitude of other individuals and information about the external factors. Individuals are socially linked by relationships mediated through a series of intertwining interactions and resulting in a highly diverse social network. This complexity can be modeled by Bayesian networks or more general Markov fields. This notion is a natural generalization of Markov chains to dynamic and spatial processes, whose domains are not necessarily linearly ordered.

6.7.1 Simplified Model: Direct and Indirect Interdependencies

In order to demonstrate the dynamics of attitude change, we begin with representing the public in groups with similar attitudes. Empirical attitude research structures the sampled population into cohorts, possibly by age, sex, income, profession, ethnicity, geographic location, political party affiliation, etc. We assume that population may be divided into “similar-attitude” groups such that an individual has a higher probability of sharing the same attitudes with others in the similar-attitude group than with individuals in other groups. Individuals communicate mainly with individuals within their group, but also with individuals in other groups. Figure 6.6 shows a simplified illustration of possible interactions between five similar-attitude groups that individuals of each group have the same attitudes. The groups are represented by nodes, and interactions are represented by arrows. Thus there is a link from group 3 to group 1, meaning that group 3 has an influence (positive or negative) on group 1. There are also links from group 1 to group 2 and from group 2 to 3. The example can be generalized to N groups, where the arrow from node i to node j indicates a link from i to j .

The direct links between nodes can be represented by the adjacency matrix shown in Table 6.1. An element of this $N \times N$ matrix indicates the position and possibly the strength of the direct dependency links from i to j .

In addition to the direct dependencies, individuals are also indirectly influenced by one another by chains of communications. If A is the adjacency matrix, then an element (i, j) of the matrix $A^2 = A \times A$ represents the number of sequential dependency paths of length 2 involving an intermediate group from i to j ; in general, A^l indicates the number of sequential dependency paths of length l with $l - 1$ intermediate groups from i to j . Thus, the entries of the matrix

$$A + A^2 + \dots + A^l$$

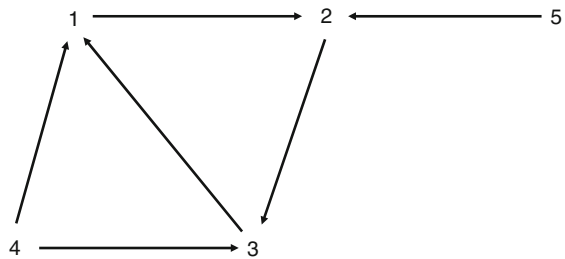


Fig. 6.6 Graph of direct relationships

Nodes	1	2	3	4	5
1	0	1	0	0	0
2	0	0	1	0	0
3	1	0	0	0	0
4	1	0	1	0	0
5	0	1	0	0	0

Table 6.1 The incidence matrix of the graph from Fig. 6.6

Fig. 6.7 Acyclic graph

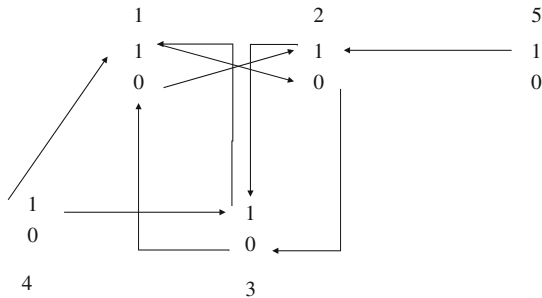
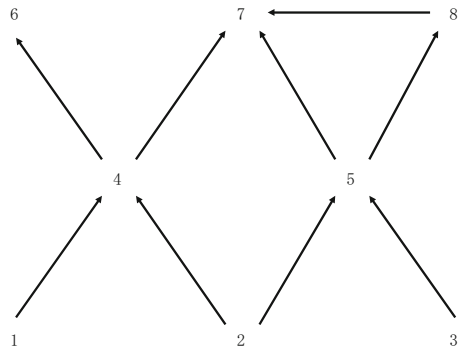


Fig. 6.8 Limit cycle attractor: waves of opinions

represent the number of all possible direct and indirect paths of length smaller or equal to l .

The graph on Fig. 6.6 has a cycle between nodes 1, 2, 3. The graph on Fig. 6.7 is acyclic which represents hierarchical structure of opinion formation.

Consider now a simple situation of how the links between individuals may affect their attitudes. Suppose that links between five groups of individuals are represented in Fig. 6.6. Figure 6.8 indicates that groups 1, 2, 3 have two possible (no–yes) attitudes $j = 0, 1$. This is represented by subnodes 0 or 1 inside of each node $i = 1, 2, 3$. We can consider 0 – 1 states of these nodes. In this example we assume the deterministic nature of interactions. Therefore assume the state $j = 1$ is settled down at nodes $i = 1, 2, 3$ at the initial time interval $t = 0$. In a general case at time $t = 0$ the state $j = 1$ is accepted only by a fraction of a group with a certain probability.

Interdependencies between groups may change their attitudes. Arrows in the graph of Fig. 6.9 indicate that individuals of the group 1 are influenced by group 3 in the sense that it expresses solidarity with group 1 taking the same opinion. Group 2 is antagonistic (in opposition) to group 1. Group 3 has the solidarity with group 2.

It easy to see a cyclic change of responses in time. When individuals of group 2 learn about the attitude of group 1 (state of node $i = 1$), the group 2 changes the attitude to the opposite. Since the initial state of all nodes $j = 1$, the next state of

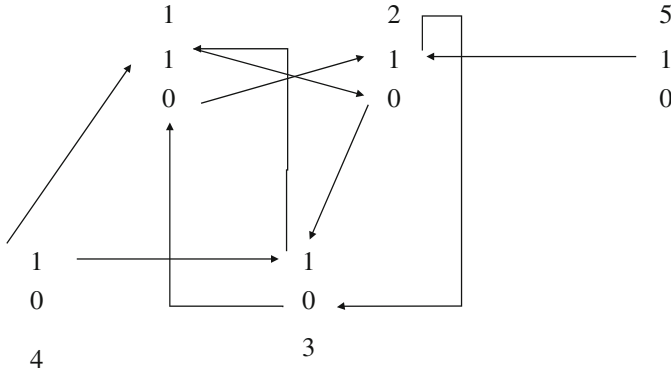


Fig. 6.9 Fixed point attractors

node 2 is $j = 0$ which triggers changes of states at nodes 3, 1 to $j = 1$. These changes lead again to state $j = 1$ at nodes 1, 2, 3, and so on.

Now suppose that group 3 is antagonistic to the group 2 (see Fig. 6.9). If group 2 learns first the response of group 1 it changes the attitude 1 to 0 and so on until attitudes reach values (1, 0, 1) for groups 1, 2, 3, respectively. If group 3 learns first the attitude of group 2, the attitudes are settled down at states (0, 1, 0). Hence, the behavior of the attitudes may display two attractors: (1, 0, 1) and (0, 1, 0).

These simple examples suggest that there may be waves of attitudes. Any opinion survey at a particular time may not represent opinion at a later time. These examples also demonstrate that delays in the learning of attitudes may change the pattern of the overall dynamics towards different attractors. In the model of the next section we take more general point of view on driving forces of attitude changes. It is assumed that members of a group may react differently at attitudes of other groups. They also may “hesitate” to react as the opposition or the solidarity.

6.7.2 Model Formulation

In the examples of the previous section individuals of each group have the same attitude. The individuals of a group also react in the same manner at “signals” (attitudes) from other groups. In this section we undertake a more general view. It is assumed that only a fraction p_j^{it} of members, the group i have attitude j at time t . Thus in the previous examples there may be fractions p_0^{it}, p_1^{it} of each group $i = 1, 2, 3$ with attitudes 0 and 1 at time t . Of course, $p_0^{it} + p_1^{it} = 1$.

Individuals of a group communicate with different individuals of other groups. Therefore, their attitudes are influenced by random samples of information from adjacent (neighbor) groups. Individuals may form their attitudes at a particular time interval t , on the basis of various rules besides just “solidarity” or “opposition”

principles. For example, they may follow the majority attitude from a sampled opinions. In general, such behavior induces a conditional probability for an individual of a group i to take an attitude from given set of possible attitudes. This probability is conditioned on their current attitudes, the attitudes of adjacent groups and some exogenous variables, or external factors.

Let us now formulate the model precisely. To do so, we must represent the driving forces of attitude changes or dependencies between groups as well as dependencies of these relations on external factors. We opted for probabilistic description based on conditional distributions.

The model distinguishes N groups $i = 1, \dots, N$ of individuals. The number of the relevant groups depends on the issue under consideration and on the level of detail represented in the model.

Individuals display different attitudes to the given issue, ranging from hostile to very favorable. We assume that there is a finite number M of possible attitudes. The attitudes of individuals of group i are described by the random variable ζ_i which takes values from 1 to M . In other words, we assume that individuals from group i statistically follow the same pattern of attitude formation given by the distribution of ζ_i . In this sense, we can say that individuals of group i share approximately the same view. The attitudes of the population are described by random vector $\zeta = (\zeta_1, \dots, \zeta_N)$. A fixed value of this vector is denoted by z and the set of all possible attitudes by Z . Let us denote by p_j^{it} the probability that a member of group i assumes the attitude j at time t :

$$p_j^{it} = \mathcal{P}(\zeta_i^t = j)$$

Naturally,

$$\sum_{j=1}^M p_j^{it} = 1, \quad p_j^{it} \geq 0$$

Interactions between individuals are represented as a graph similar to that described in Sect. 6.7.1. Groups $i = 1, \dots, N$ correspond to nodes of the graph and direct links between individuals of groups are represented by arrows between nodes. Thus there are two sets: nodes $V = 1, \dots, N$ and the set of arrows (directed arcs) U . Let us denote this as $G = (V, U)$. If nodes i, j belong to V , $i, j \in V$ and there is an arrow from i to j , $(i, j) \in U$ then i is an adjacent to j node. Define as V_j the set of all nodes adjacent to the j node. Let z_{V_j} be a subvector of the vector of attitudes indexed by V_j . For example, in the case of dependencies indicated by graph in Fig. 6.6 we have $V_1 = \{1, 3, 4\}$, $V_2 = \{1, 2, 5\}$, $V_3 = \{2, 3, 4\}$, $V_4 = \{\emptyset\}$, $V_5 = \{\emptyset\}$ where \emptyset is the symbol of the empty set. Then $z_{V_1} = (z_1, z_3, z_4)$, $z_{V_2} = (z_1, z_2, z_5)$.

Individuals change their attitudes depending on current attitudes of their own and adjacent groups, and on a vector x of exogenous external factors or variables. The attitude formation is described by a conditional distribution $H^i(z_i | z_{V_i}, x)$ for each individual of group i : the probability for an individual of group i to have an attitude $z_i = 1, 2, \dots, M$ when attitudes of groups V_i are Z_{V_i} and external factors have values x .

We assume that function H^i can be derived on the basis of appropriate questionnaires answered by a representative member of individuals from each group, such as: “What is your attitude (from 1 to M) if attitudes of the adjacent groups are z_{V_i} and the environment is $x = (x_1, \dots, x_n)$?”. Functions H^i may also depend on time interval t , but we skip it in order to simplify the notation. Functions H^i define the dynamics of the attitude change according to the following relation:

$$p_j^{i,t+1} = \sum_{z_{V_i} \in Z} H^i(\zeta_i^{t+1} = j | \zeta_{V_i}^t = z_{V_i}; x^t) \mathcal{P}(\zeta_{V_i}^t = z_{V_i}) \quad (6.26)$$

The groups i with $V_i = i$ can be identified with “leaders”, which influence opinions of other individuals but are not influenced themselves.

To define completely the dynamics of the system described by (6.26) it is necessary to fix initial attitudes distributions for $t = 0$. In case when the corresponding graph of direct influences is acyclic, it is enough to define these distributions for nodes i such that $V_i = i$.

Equation (6.26) together with initial distributions allows us in principle to calculate $p_j^{i,t}$ for any $t \geq 0$. Of course, for complex graphs it is practically impossible to derive analytical formulas for $p_j^{i,t}$ as functions of external factors x . The existence of analytical expressions for all $p_j^{i,t}$ provides an easy tool to analyze implications of changes in x . The next section is devoted to the analysis of attitudes changes in the case when such a possibility does not exist. The approach is based on the stochastic version of (6.26) dealing directly with random variables ζ_i^t , $i = 1, \dots, N$ by using the Monte-Carlo simulation techniques. Instead of H^i , $i = 1, \dots, N$ the approach allows also to use myopic rules for generating random changes of ζ^t , i.e., the approach allows to analyze cases when functions H^i are given implicitly. One such important case arises in the situations when individuals form their attitudes by asking acquaintances from adjacent groups and use some simple rules based on majority or minority of sampled attitudes. Let us now formulate some problems that are important in this context.

Problem 1. Evaluation of attitudes

The objective here is to predict attitudes of various population groups. As we have seen in the previous section, the attitudes of different groups change in intricate ways and are subject to changes in external factors, as well as direct and indirect dependencies. Direct dependencies involve relatively few adjacent groups, while indirect dependencies and external factors may involve all or almost all population groups. Thus, in Fig. 6.1 group 2 is affected directly only by groups 1, 5. But there exists a path from 3 to 2, and from 4 to 2. Therefore, indirectly individuals of group 2 are affected by all groups. The direct dependencies are much easier to study experimentally through surveys and questionnaires. Suppose that we managed to study the direct dependencies between attitudes of different adjacent population groups. The problem is to predict the public attitudes as the result of complex direct and indirect interactions by using the information about direct dependencies. As was outlined in

the previous section, this problem involves the calculation of all possible direct and indirect paths by using the adjacency matrix (Table 6.1). Formally the problem is formulated as follows. Given conditional distributions $H^i(z_i|z_{V_i}; x)$ for $i = 1 : N$ and the values of x^t for $t = 0, 1, \dots, T - 1$ find distributions $\mathcal{P}(\xi_i^T)$, $i = 1 : N$ of random variables ξ_i^T , i.e., the public attitudes at time T .

Problem 2. Response interpretation

This problem deals with the interpretation (identification) of public response to a mixture of events and efforts which influence the public. This interpretation is made on the basis of our knowledge which has two components. The first component is the a priori information on direct dependencies deduced from responses in the past. The second component consists of new direct observations of attitudes for some groups to a given new issue. This type of knowledge can be called a posteriori knowledge. The response interpretation deals with the following questions:

- Suppose that we have direct observations on the attitudes of only some selected groups, or we have observations of aggregated response from several groups. How can we recover attitudes of unobserved groups?
- How to use the newly acquired a posteriori knowledge to update our knowledge about direct dependencies between groups?
- Often a public response is the result of mixture of different, sometimes conflicting events and efforts. What is the contribution of each single event to the attitude dynamics?

Formally these problems can be formulated as follows. Let us consider only first problem. Denote by $V_{\mathcal{E}}$ the set of observed groups. Given conditional distributions $H^i(z_i|z_{V_i}; x)$ for $i = 1 : N$, the values of x^t and distributions $\mathcal{P}(\xi_i^t)$, $i \in V_{\mathcal{E}}$ for $t = 0, 1, \dots, T - 1$ find distributions $\mathcal{P}(\xi_i^t)$ for $i \notin V_{\mathcal{E}}$.

Problem 3. Sensitivity analysis

Here we want to analyze a sensitivity of attitudes with respect to changes of environmental variable x . Through such an analysis we may find that attitudes are especially resistant to changes in certain directions or in certain positions. For example, in siting a waste processing facility, it is necessary to analyze a choice of its size, decide on a distance between facility and population centers, choose routes of the waste transportation, etc. Different population groups react differently on different options. Small changes in critical parameters may considerably affect the public attitudes, while substantial and possibly costly changes in non-critical parameters will not change the public response. The objective of the sensitivity analysis is to identify the critical parameters utilizing the knowledge of the direct dependencies and how these dependencies are affected by changes in environmental parameters.

In terms of our model the sensitivities of public responses is defined in terms of changes in response distribution $\mathcal{P}(\cdot)$ with respect to parameters x . This leads to the following formulation.

Given conditional distributions $H^i(z_i|z_{V_i}; x)$ for $i = 1 : N$ and the values of x^t for $t = 0, 1, \dots, T - 1$ estimate derivatives of distributions $\mathcal{P}(\xi_i^T), i = 1 : N$ with respect to x^0, \dots, x^T .

Note that distributions $\mathcal{P}(\cdot)$ depend on x indirectly through conditional distributions H , what leads to a challenging problems analyzed in next sections.

Problem 4. Social learning

As it is emphasized in social psychology people receive information from their social environment. A lack of connectivity between them develops clusters of people sharing similar views in a more heterogeneous population. Traditionally, it is assumed that a change in thought or behavior is only a reaction to some external factors or stimulus. The proposed model emphasizes the existence of intrinsic changes generated by interdependencies between individuals in the absence of any external factor. This dynamics may be perturbed by external factors activated at different times leading to more or less complex patterns of alternative dynamics. How can we learn the variety of alternative attitudinal developments and how we can characterize them? What are optimistic and pessimistic “scenarios” of such developments? Can attitudes reach critical levels? Answers to such questions depend not only on existing links between individuals but also on paths of activated external factors. The main problem is to use a model in order to learn possible alternative scenarios and their outcomes. For example, in the debates on siting a waste processing facility, there is a possibility to change sizes of facilities, their locations, premiums and other compensations in order to change public responses.

The power of a model is its ability to learn patterns of possible responses of the system without time consuming real observations and trial-and-error experiments. In our case the proposed model supports identification of paths of external factors leading to different outcomes, for example either decreasing a social tension, or achieving the worst case situation. In order to conduct such analysis we need to introduce a set of “performance” indicators or “score” functions distinguishing one trajectory of attitudes from another. For example, if c_{ij} is relative importance of attitude j by individuals of group i , then the cumulative score of a trajectory of attitudes can be expressed by the following score (performance) function:

$$F(x) = \sum_{t=0}^T \sum_{i=1}^N \sum_{j=1}^M c_{ij} p_j^{it}, \quad (6.27)$$

which implicitly depends on external factors x through conditional probabilities in (6.26).

The sensitivity analysis can indicate changes in x which lead to increasing value of an indicator $F(x)$. Using this information, it is possible to identify, for example, the worst or the best case sets of possible external factors.

The fundamental complexity of such type of problems is due to the probabilistic nature of $F(x)$ and implicit dependencies on variables x . Next sections discuss tools enabling to deal with the involved complexity.

6.7.3 Bayesian Networks and Markov Fields

The Bayesian net is a powerful tool which was developed primarily to deal with stochastic problems defined on acyclic graphs (see Fig. 6.8) We show that algorithms developed for the Bayesian nets constitute the building blocks for more general algorithms which can deal also with general graphs.

Bayesian nets are specifically designed for cases when the vector of random parameters ζ can have considerable dimension and/or it is difficult to come up with traditional parametric model of the joint distribution of random parameters.

The cause–effect structure is associated with the vector of random parameters $\zeta = (\zeta_1, \dots, \zeta_i, \dots, \zeta_N)$. That is, for any parameter ζ_i the set of indices V_i is selected such that the elements of subvector $\{\zeta_j\}_{j \in V_i}$ can be identified with “causes” of ζ_i . Vector ζ_i is changed in time leading to a random path or trajectory $\zeta^t, t = 0, 1, \dots$. In the proposed model, the groups of a population, say in a given region, are represented as nodes of a graph. Direct dependencies between groups are represented by arrows indicating directions of communications. Node i of the graph has different random states ζ_i^t at time t reflecting different attitudes of the group. Thus the stochastic dynamics of attitudes is characterized by a random vector ζ^t with dependent components. The important feature is that changes in a component ζ_i^t are triggered only by its “neighbors” $\zeta_j^t, j \in V_i$, apart from changes in external factors.

Such stochastic processes define Markov random fields. They can be regarded as a generalization of Markov processes to situations when the cause effect structure is not necessarily linearly ordered. Let us start the formal exposition by gathering the basic definitions which will be used in next sections. Consider a directed graph defined as a pair of two sets (V, U) : nodes V and connecting them arrows (directed arcs) U . An example of such graph is given in Fig. 6.6 with $V = \{1, 2, \dots, 9\}$. For each node $v \in V$ let us define the set of parents

$$c(v) = \{j | (j, v) \in E\}, \tag{6.28}$$

and the set of descendants $d(v)$ as the set of all nodes \bar{v} from V for which there exists directed path which originates in v and terminates in \bar{v} .

Consider a set of random variables $\zeta_V = \{\zeta_v, v \in V\}$ indexed by nodes from V and defined on appropriate probability space $(\Omega, \mathbb{B}, \mathbb{P})$. Suppose that W is an arbitrary subset of V by $v \setminus \{v \cup d(v)\}$. A Markov random field ζ is characterized by the property

$$\mathbb{P}(\zeta_v | \zeta_{W \cup c(v)}) = \mathbb{P}(\zeta_v | \zeta_{c(v)}) \tag{6.29}$$

If G is a directed acyclic graph, then (ζ, G) which satisfies (6.29) is called *Bayesian network*.

Suppose now that $H(\zeta)$ is the joint distribution of ζ . Then it follows from the definition of the Bayesian network that

$$H(\zeta) = \prod_{v \in V} H^v(\zeta_v | \zeta_{c(v)}) \quad (6.30)$$

where $H^v(\zeta_v | \zeta_{c(v)})$ is a conditional distribution function of ζ_v given $\zeta_{c(v)}$.

Coming back to the model of Sect. 6.3 we see that it fits in the framework of Bayesian networks and Markov fields. The essential new feature is the dependence of the conditional distribution functions H^v on the vector of external variables x from a feasible set $X \subseteq R^n$:

$$H^v(\cdot) = H^v(\zeta_v | \zeta_{c(v)}, x) \quad (6.31)$$

The change of $x \in X$ affects the interactions between groups.

6.7.4 Sensitivity Analysis

The analysis aims at examining attitudes with respect to changes in external factors x . These changes are characterized by certain indicators or “score” functions such as (6.27) is a function of ζ and possibly x , e.g., $f(x, \zeta)$.

Several questions can now be considered. For example, how can we estimate the expected outcome

$$F(x) = \mathbb{E}f(x, \zeta) \quad (6.32)$$

at some point x ? How sensitive is this value with respect to changes of parameters x ? What are the most critical parameters? What is the value of the performance indicator at point $x + \delta x$ where δx is a small perturbation of x ? These questions are not the trivial because each estimation of the value of $F(x)$ can be very time consuming taking into account indirect interdependencies of the network.

More precisely, in order to evaluate the sensitivity of $F(x)$ we need to develop algorithms for estimating the value of the gradient of this indicator. That is, for a given x we need to compute vector ξ such that

$$\mathbb{E}\xi = F_x(x) = \frac{d}{dx} \mathbb{E}f(x, \zeta)$$

One possibility is to use the finite differences:

$$\xi = \sum_{i=1}^n \frac{\hat{F}(x + \Delta e_i) - \hat{F}(x)}{\Delta} e_i,$$

where $\hat{F}(x)$ is an estimate of the value of function $F(x)$ at point x . In this case, however, it is necessary to compute at least $n + 1$ estimates of the performance measure which may be too demanding computationally.

Let us introduce simple but useful differentiation formulas for the gradient of function $F(x)$, which are used in stochastic optimization.

Theorem 6.1. *Suppose that*

1. *Random variables ζ_v have conditional densities $h(\zeta_v|\zeta_{c(v)}, x)$*
2. *Functions $h(\zeta_v|\zeta_{c(v)}, x)$, $f(x, \zeta)$ are differentiable with respect to x uniformly with respect to ζ . Then $F(x)$ defined in (6.32) is differentiable and*

$$F_x(x) = \mathbb{E} \left\{ f_x(x, \zeta) + f(x, \zeta) \sum_{v \in V} \frac{h_x(\zeta_v|\zeta_{c(v)}, x)}{h(\zeta_v|\zeta_{c(v)}, x)} \right\} \quad (6.33)$$

Proof.

From (6.30) we have the following expression for $F(x)$:

$$F(x) = \int f(x, \zeta) \prod_{v \in V} h(\zeta_v|\zeta_{c(v)}, x) \prod_{v \in V} d\zeta_v \quad (6.34)$$

Under assumptions of the theorem we can change the order of differentiation and integration, which yields:

$$\begin{aligned} F_x(x) &= \int \frac{d}{dx} \left(f(x, \zeta) \prod_{v \in V} h(\zeta_v|\zeta_{c(v)}, x) \right) \prod_{v \in V} d\zeta_v \\ &= \int f_x(x, \zeta) \prod_{v \in V} h(\zeta_v|\zeta_{c(v)}, x) \prod_{v \in V} d\zeta_v \\ &\quad + \sum_{w \in V} \int f(x, \zeta) h_x(\zeta_w|\zeta_{c(w)}, x) \prod_{v \in V, v \neq w} h(\zeta_v|\zeta_{c(v)}, x) \prod_{v \in V} d\zeta_v \\ &= \int \left(f_x(x, \zeta) + f(x, \zeta) \sum_{v \in V} \frac{h_x(\zeta_v|\zeta_{c(v)}, x)}{h(\zeta_v|\zeta_{c(v)}, x)} \right) \prod_{v \in V} h(\zeta_v|\zeta_{c(v)}, x) \prod_{v \in V} d\zeta_v \end{aligned}$$

The proof is completed. \square

This theorem can also be proved by taking the logarithm from the expression under the integral in (6.35) and differentiating it. The theorem is similar to the Theorem 6.1 from [15].

Similar result holds when each of the random variables ζ_v takes finite number of values and instead of conditional densities we have conditional probabilities $P(\zeta_v|\zeta_{c(v)}, x)$. If these probabilities are differentiable with respect to x we obtain the following expression for $F_x(x)$:

$$F_x(x) = \mathbb{E} \left\{ f_x(x, \zeta) + f(x, \zeta) \sum_{v \in V} \frac{P_x(\zeta_v|\zeta_{c(v)}, x)}{P(\zeta_v|\zeta_{c(v)}, x)} \right\} \quad (6.35)$$

Note that the second terms in expressions (6.33) and (6.35) can be interpreted as sums of likelihood ratios [17, 27]. The calculation of exact values $F_x(x)$ is possible only in exceptional cases for simple networks.

Let us now present estimation algorithms which exploit the structure of Bayesian network and expressions (6.33) and (6.35) in order to obtain statistical estimates of $F(x)$ and $F_x(x)$. We shall make reference to discrete case and use (6.35); the continuous case is treated similarly. The simplest estimation scheme is the following:

$$\xi = \frac{1}{K} \sum_{k=1}^K \left(f_x(x, \zeta^k) + f(x, \zeta^k) \sum_{v \in V} \frac{P_x(\zeta_v^k | \zeta_{c(v)}^k, x)}{P(\zeta_v^k | \zeta_{c(v)}^k, x)} \right) \quad (6.36)$$

where $\zeta^k, \zeta_v^k, \zeta_{c(v)}^k, k = 1 : K, K \geq 1$ are independent observations of random vectors $\zeta, \zeta_v, \zeta_{c(v)}$ respectively. Stochastic vector ξ is termed as stochastic gradient. The estimation scheme (6.36) in the context of Bayesian networks is called also as Logic Sampling. The estimator (6.36) requires $K \geq 1$ observations (“scenarios”) of attitudes ζ . This scenarios can be sequentially generated by using the Monte-Carlo simulation techniques. An arbitrary scenario generating cycle $k = 1 : K$ have the following simple steps:

1. *Initialization.* Select a node $v \in V$ such that $c(v)$ is the empty set, $c(v) = \emptyset$. Since the graph G is acyclic such nodes exist and we interpreted them as sources of opinions or leader nodes. Sample the component ζ_v from unconditional distribution $P(\zeta_v | \zeta_{c(v)}, x)$ and denote the result as ζ_v^k . Perform this step for all nodes with empty parent set $c(v)$.
2. *Selection.* Select a not yet sampled node $v \in V$ such that all random variables $\zeta_j, j \in c(v)$ have been sampled already during current scenario generating cycle. Suppose that the observation of ζ_j is $\zeta_j^k, j \in c(v)$.
3. *Sampling.* Sample random variable ζ_v and obtain an observation ζ_v^k . Steps 2 and 3 are repeated until all random components of ζ are sampled. In other words, until all groups reveal their opinions.

Note that by using observations $\zeta^k, k = 1 : K, K \geq 1$ along with the estimates of the gradient F_x we obtain the estimate of $F(x)$ itself. For example, $F(x)$ may correspond to the probability for ζ to belong to a certain desirable domain [for example, majority of attitudes 1 (yes) to the competing attitudes 0 (no)] we obtain the estimates of this probability and its sensitivity ξ .

There may be other sampling schemes known as Evidence Weighting [4] and Gibbs Sampling [16] which are particularly useful for the solution of the response interpretation problem (Problem 2 from Sect.6.7.2). Such schemes use the structure of the network and a posteriori distributions instead of $P(\zeta_v | \zeta_{c(v)}, x)$. The estimates of stochastic gradient obtained above can be used for obtaining the optimal values of parameters using stochastic quasigradient methods like in [7–9].

6.7.5 General Interdependencies

Let us extend the analysis of the previous section to the case of cyclic graphs. The exposition follows closely the exposition of the previous section and therefore we concentrate on the new features only. The most important among these features is

the dynamic aspect of attitudes change which in the case of conventional Bayesian network was reduced to analysis of their propagation from “leaders” to other groups.

Let us fix the time horizon $[0, l]$ during which we study the attitude dynamics and sensitivity estimates. Such estimates will be called l -links estimates. Generally, the variables x may change during this time period: $x = (x^0, \dots, x^{l-1})$. For the sake of simplicity we shall derive our estimates for the case when $x^0 = x^1 = \dots = x^{l-1}$ denoting this constant vector by x . The general case does not bring any conceptual difficulties and is treated similarly.

Consider again the case when conditional distributions have densities. Consider the following performance measure which is a generalization of the measure (6.34):

$$F(x) = \int f(x, \bar{\zeta}^l) h(x, \bar{\zeta}^l) d\bar{\zeta}^l \tag{6.37}$$

where for $t = 0, \dots, l$ we denoted by $h(x, \bar{\zeta}^t)$ the joint density of the random vector $\bar{\zeta}^t = (\zeta^0, \dots, \zeta^t)$ with $\zeta^t = (\zeta_1^t, \dots, \zeta_N^t)$ and

$$d\bar{\zeta}^l = \prod_{t=1}^l \prod_{v \in V} d\zeta_v^t$$

The density of $\bar{\zeta}^t$ is connected with the density of $\bar{\zeta}^{t-1}$ for $t = 1, \dots, l$ with the following relation:

$$h(x, \bar{\zeta}^t) = h(x, \bar{\zeta}^{t-1}) \prod_{v \in V} h(\zeta_v^t | \zeta_{c(v)}^{t-1}, x) \tag{6.38}$$

with the initial distribution $h(x, \bar{\zeta}^0)$ density being simply

$$h(x, \bar{\zeta}^0) = \prod_{v \in V} h(\zeta_v^0) \tag{6.39}$$

Combining expressions (6.37)–(6.39) we obtain the basic formula for the performance measure:

$$F(x) = \int f(x, \bar{\zeta}^l) \prod_{v \in V} h(\zeta_v^0) \prod_{t=1}^l \prod_{v \in V} h(\zeta_v^t | \zeta_{c(v)}^{t-1}, x) \prod_{t=1}^l \prod_{v \in V} d\zeta_v^t \tag{6.40}$$

Utilizing this expression similarly to the previous section we obtain the following result:

Theorem 6.2. *Suppose that*

1. *Random variables ζ_v^t have conditional densities $h(\zeta_v^t | \zeta_{c(v)}^{t-1}, x)$ for $t = 1, \dots, l$ and densities $h(\zeta^0)$ for $t = 0$.*

2. Functions $h(x, \zeta_v^t | \zeta_{c(v)}^t)$, $f(x, \bar{\zeta}^l)$ are differentiable with respect to x uniformly with respect to $\bar{\zeta}^l$. Then $F(x)$ defined in (6.37) is differentiable and

$$F_x(x) = \mathbb{E} \left\{ f_x(x, \bar{\zeta}^l) + f(x, \bar{\zeta}^l) \sum_{t=1}^l \sum_{v \in V} \frac{h_x(\zeta_v^t | \zeta_{c(v)}^{t-1}, x)}{h(\zeta_v^t | \zeta_{c(v)}^{t-1}, x)} \right\} \quad (6.41)$$

As in the previous section similar result holds when each of the random variables ζ_v^t takes a finite number of values, and instead of conditional densities we have conditional probabilities $P(x, \zeta_v^t | \zeta_{c(v)}^t, x)$. If these probabilities are differentiable with respect to x , then we obtain the following expression for $F_x(x)$:

$$F_x(x) = \mathbb{E} \left\{ f_x(x, \bar{\zeta}^l) + f(x, \bar{\zeta}^l) \sum_{t=1}^l \sum_{v \in V} \frac{P_x(\zeta_v^t | \zeta_{c(v)}^{t-1}, x)}{P(\zeta_v^t | \zeta_{c(v)}^{t-1}, x)} \right\} \quad (6.42)$$

The l -stage sensitivity estimate based on the logic sampling in the discrete case takes the form:

$$\xi = \frac{1}{K} \sum_{k=1}^K \left(f_x(x, \bar{\zeta}^{kl}) + f(x, \bar{\zeta}^{kl}) \sum_{t=1}^l \sum_{v \in V} \frac{P_x(\zeta_v^{kt} | \zeta_{c(v)}^{k,t-1}, x)}{P(x, \zeta_v^{kt} | \zeta_{c(v)}^{k,t-1}, x)} \right) \quad (6.43)$$

where $\bar{\zeta}^{kt}, \zeta_v^{kt}, \zeta_{c(v)}^{kt}$, $t = 0, 1, \dots, l$, $k = 1 : K$, $K \geq 1$ are independent observations of random vectors $\bar{\zeta}^t, \zeta_v^t, \zeta_{c(v)}^t$ respectively. The vector ξ defines a stochastic gradient of the function $F(x)$ in (6.37).

Observations ζ^{kt} , $k = 1 : K$ of random vectors ζ^t , $t = 0, 1, \dots, l$ can be simulated similar to acyclic graphs of the previous section. In fact, the study of l -stage interdependencies between groups (nodes) on general graphs can be reduced to the study of l -stage sensitivity estimates on acyclic graphs.

This reduction of l -stage sensitivity analysis to acyclic graphs provides a general approach to the study of the general nets with cyclic graphs. Keeping in mind this possibility let us describe the Monte-Carlo simulation procedure for the generation $k = 1 : K$ scenarios ζ , $t = 0, 1, \dots, l$ describing l -stage propagation effects of public attitudes:

1. *Initialization.* Leader nodes of the equivalent acyclic graph (see Fig. 6.6) correspond to $t = 0$. Therefore, for all $v \in V$ sample the values ζ^{kt} of random variables ζ_v^t , $t = 0$.
2. *Selection.* Select all nodes $v \in V$ sequentially for $t = 1 : K$.
3. *Sampling.* For a sampled node v at step t sample random variables ζ_v^{kt+1} from distribution $P(\zeta_v^{t+1} | \zeta_v^t = \zeta_v^{kt}, x)$. Steps 2, 3 are repeated until all components ζ^{kt} of vectors ζ^t , $t = 0, 1, \dots, l$ are sampled.

Scenarios ζ^{kt} , $t = 0, 1, \dots, l$, $k = 1 : K$ of public attitudes allow to estimate vector $F_x(x)$ according to (6.43). As in the previous section this vector indicates directions of changes in x towards the increase of indicator $F(x)$.

6.8 Conclusion

In this paper we set the stage for further development of modeling and decision support tools for analysis of design and deployment of robust information infrastructure using as the case study one component of this infrastructure: advanced mobile data services. The methodologies utilized here, especially the portfolio theory and the Bayesian nets combined with the stochastic optimization, have a potential to be exploited also for solutions of similar problems arising in other types of infrastructure.

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Part III
**Analysis and Optimization of Technical
Systems and Structures Under Uncertainty**

Chapter 7

Optimal Ellipsoidal Estimates of Uncertain Systems: An Overview and New Results

F.L. Chernousko

Abstract The method of ellipsoids for the guaranteed state estimation of uncertain dynamical systems is associated with optimal two-sided ellipsoidal bounds for reachable sets of the systems. Being based on the set-membership approach to uncertainties, the method can be regarded as a natural counterpart to well-known stochastic, or probabilistic, techniques. Basic concepts and results of the method are outlined, and certain results are presented. Various possible applications to problems in control, estimation, and observation are considered.

7.1 Introduction

Dynamical systems subjected to unknown but bounded perturbations appear in numerous applications. The set-membership approach that is a natural counterpart to the well-known stochastic, or probabilistic, one makes it possible to obtain guaranteed estimates on reachable sets and thus to evaluate the family of all possible trajectories of the perturbed system.

In the framework of the set-membership approach, the ellipsoidal estimation seems to be the most efficient technique. Among its advantages are the explicit form of approximations, invariance with respect to linear transformations, possibility of optimization, etc. The earlier results on the ellipsoidal estimation were presented in [33]. The concept of optimality for two-sided (inner and outer) approximating ellipsoids was first introduced in [1] and generalized, extended, and summarized in books [2, 3, 5].

In this paper, basic concepts and results of the method of optimal ellipsoids are outlined, and certain recent results are presented.

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Dynamical systems subjected to bounded controls and/or perturbations are considered. For these systems, nonlinear differential equations are obtained that describe the evolution of the optimal ellipsoids representing two-sided (inner and outer) estimates for reachable sets. The approximating ellipsoids depend on the choice of the optimality criterion (e.g., volume of ellipsoids, sum of their squared axes), and on the notion of local/global optimality.

Various useful properties of the optimal approximating ellipsoids have been established. Asymptotic behavior of the ellipsoids near the initial point and at infinity have been studied. As a rule, the nonlinear equations for these ellipsoids are to be integrated numerically. However, certain explicit analytical solutions have been obtained.

Outer and inner ellipsoidal approximations can be used for various applications in control and estimation, including two-sided approximations for optimal control and differential games, analysis of practical stability and parameter excitation, state estimation in the presence of observation errors, control in the presence of uncertain perturbations, etc.

7.2 Reachable Sets

Consider a dynamical system subjected to control or disturbance and described by a system of ordinary differential equations

$$\dot{x} = f(x, u, t), \quad t \geq s \quad (7.1)$$

Here, $x = (x_1, \dots, x_n)$ is the vector of state, t is time, s is the initial time instant, $u = (u_1, \dots, u_m)$ is the vector of control or disturbance, and f is a given function. At each time instant, vector $u(t)$ should belong to the given set $U(t)$, so that

$$u(t) \in U(t) \subset R^m, \quad t \geq s. \quad (7.2)$$

The initial state belongs to a given initial set:

$$x(s) \in M \subset R^n. \quad (7.3)$$

An alternative description of the system (7.1)–(7.3) is given by the differential inclusion:

$$\dot{x} \in f(x, U(t), t), \quad x(s) \in M.$$

For systems under consideration, the notion of a reachable set is introduced.

The *reachable*, or *attainable*, set $D(t, s, M)$ of system (7.1) for $t \geq s$ is defined as the set of all end points $x(t)$ at the instant t of all state trajectories $x(\cdot)$ compatible

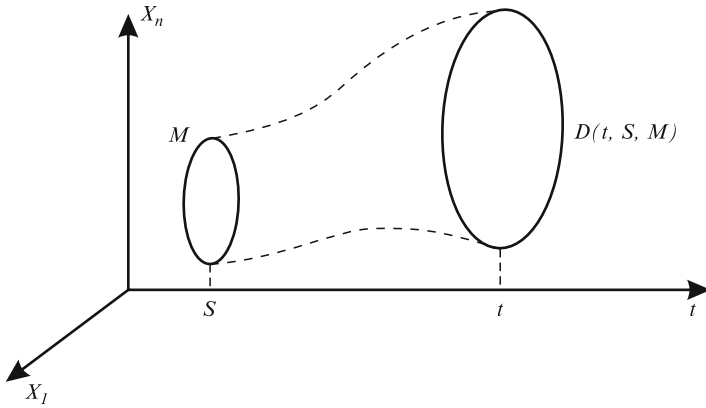


Fig. 7.1 Reachable set

with (7.1)–(7.3). The reachable set has the following evolutionary property

$$D(t, s, M) = D(t, \tau, D(\tau, s, M)) \tag{7.4}$$

that holds for all $\tau \in [s, t]$. Reachable sets are important for systems subjected to control or disturbance because they give a description of all possible states of the system at any given time t (Fig. 7.1).

In various applications, it is often required to verify whether a given state x^* is reachable at the given instant of time t^* , i.e., if the inclusion

$$x^* \in D(t^*, s, M) \tag{7.5}$$

holds. However, the practical determination of reachable sets in the n -dimensional space presents usually serious computational difficulties. In a number of cases, it is sufficient to obtain simple and efficient two-sided (inner and outer) bounds $D^-(t)$ and $D^+(t)$ on reachable sets such that the following inclusions

$$D^-(t) \subset D(t, s, M) \subset D^+(t) \tag{7.6}$$

are true for $t \geq s$. If the bounds D^- and D^+ are known, then the inclusions

$$x^* \in D^-(t^*), \quad x^* \in D^+(t^*) \tag{7.7}$$

can serve, respectively, as a sufficient and necessary conditions for the inclusion (7.5).

If we deal with a control system and u in (7.1) is a control to be chosen, then the inner bound $D^-(t)$ is important. The first inclusion (7.7) implies that the state x^* is

reachable at the instant t^* , and there exists an admissible control $u(t)$ bringing the system to this state.

For an uncertain system, $u(t)$ is a disturbance, and the outer bound $D^+(t)$ provides a guaranteed estimate for all possible trajectories: the system cannot reach any state outside the set $D^+(t^*)$ at the instant t^* .

In the well-known stochastic (probabilistic) approach to uncertain systems, the Gaussian distribution plays a specific role and results in the simplest mathematical description. To some extent, ellipsoidal sets for the set-membership approach to uncertainties are analogous to the Gaussian distributions. In fact, the sets of constant probability for the n -dimensional Gaussian distribution are the surfaces of n -dimensional ellipsoids. Moreover, there is a certain similarity between stochastic systems subjected to the white noise and dynamical systems subjected to uncertain disturbances bounded by ellipsoids.

Denote by $E(a, Q)$ the following n -dimensional ellipsoid

$$E(a, Q) = \{x : (Q^{-1}(x - a), (x - a)) \leq 1\}, \quad (7.8)$$

where $a \in R^n$ is its center, Q is a symmetric positive definite $n \times n$ matrix, and (\cdot, \cdot) denotes the scalar product of vectors.

Ellipsoids have a number of advantages as approximating sets. They provide a satisfactory approximation for a wide class of convex sets [1–3]. For any n -dimensional convex set D , there exists an ellipsoid $E(a, Q)$ such that

$$E(a, Q) \subset D \subset E(a, n^2 Q).$$

If the convex set is symmetric with respect to some point, then there exists an ellipsoid $E(a, Q)$ for which the inclusions.

$$E(a, Q) \subset D \subset E(a, nQ)$$

hold. These inclusions provide two-sided estimates for possible approximations of any convex set by means of ellipsoids.

The class of ellipsoids is invariant with respect to affine transformations. The class of parallelepipeds has the same invariance property but ellipsoids are defined by less number of parameters: $n + n(n + 1)/2$ for ellipsoids, $n(n + 1)$ for parallelepipeds. Note that rectangular parallelepipeds are defined by the same number of parameters as ellipsoids but they are not invariant with respect to affine transformations.

The last but not the least advantage of ellipsoids is the following one. Simple explicit formulas were obtained [1–3] for the basic algebraic operations for ellipsoidal sets, namely, for the multiplication by a constant, the sum, and intersection of ellipsoids. These operations are, in a certain sense, optimal and provide a basis for the ellipsoidal estimation of dynamical systems subjected to controls and/or uncertain disturbances.

The ellipsoidal state estimation occurs to be the most effective technique in the framework of the set-membership approach. In what follows, we will remind the basic concepts and results of this technique.

7.3 Ellipsoidal Bounds

Let us specify the general system (7.1) and consider a linear system of ordinary differential equations

$$\dot{x} = A(t)x + B(t)u + f(t), \quad t \geq s. \quad (7.9)$$

Here, $x \in R^n$ is the n -vector of state, $u \in R^m$ is the m -vector of control or unknown disturbances, A is an $n \times n$ matrix, B is an $n \times m$ matrix, and f is an n -vector. The matrices $A(t)$ and $B(t)$ as well as the vector $f(t)$ are given functions of time t for $t \geq s$.

Suppose the set $U(t)$ that bounds vector $u(t)$ in (7.2) is an ellipsoid. We have

$$u(t) \in E(0, G(t)), \quad t \geq s, \quad (7.10)$$

where $G(t)$ is a positive definite $m \times m$ matrix specified for $t \geq s$.

The initial set M in (7.3) is also supposed to be an ellipsoid. We have the following bound on the initial state:

$$x(s) \in M = E(a_0, Q_0), \quad (7.11)$$

where a_0 is a given n -vector, and Q_0 is a given positive definite $n \times n$ matrix.

Our goal is to obtain two-sided bounds (7.6) on reachable sets, where the bounding sets $D^-(t)$ and $D^+(t)$ are n -dimensional ellipsoids. In other words, we look for two families of n -dimensional ellipsoids:

$$E^-(t) = E(a^-(t), Q^-(t)), \quad E^+(t) = E(a^+(t), Q^+(t)) \quad (7.12)$$

defined, according to notation (7.8), by their centers $a^-(t)$, $a^+(t)$ and positive definite $n \times n$ matrices $Q^-(t)$, $Q^+(t)$ for $t \geq s$ and such that the following inclusions hold:

$$E^-(t) \subset D(t, s, M) \subset E^+(t), \quad t \geq s. \quad (7.13)$$

In addition, we require that the families of ellipsoids $E^-(t)$ and $E^+(t)$ possess the properties of subreachability and superreachability, respectively.

The families of ellipsoids $E^-(t)$ and $E^+(t)$ defined by (7.11) are called *sub-reachable* and *superreachable*, respectively, if

$$E^-(t) \subset D(t, \tau, E^-(\tau)), \quad E^+(t) \supset D(t, \tau, E^+(\tau))$$

for all $\tau \in [s, t]$. These properties are similar to the evolutionary property (7.4) of reachable sets. It occurs that subreachable and superreachable ellipsoids have certain advantages: they can be determined efficiently in a straightforward way and provide a sufficiently good approximations of reachable sets.

7.4 Optimality

To make the approximating ellipsoids (7.12) closer to reachable sets, it is quite natural to impose certain optimality conditions upon these ellipsoids. The optimality criteria should, to some extent, reflect the “size” of ellipsoids, and the inner ellipsoid E^- should be “larger”, whereas the outer ellipsoid E^+ “smaller”, in the sense of the chosen criterion.

Let us characterize an ellipsoid $E(a, Q)$ by a scalar optimality criterion J which is a given function $L(Q)$ of the matrix Q , i.e., $J = L(Q)$. The function $L(Q)$ is defined for all symmetric positive definite matrices Q , is smooth and monotone. The monotonicity means that $L(Q_1) \geq L(Q_2)$, if $Q_1 - Q_2$ is a nonnegative definite matrix.

Consider some important particular cases of the general criterion $L(Q)$:

1. The volume of an ellipsoid [1–3] is given by

$$J = c_n (\det Q)^{1/2}, \quad (7.14)$$

where c_n is a constant depending on n .

2. The sum of the squared semiaxes of an ellipsoid is equal to

$$J = \text{Tr} Q.$$

3. A linear optimality criterion [3, 5]

$$J = \text{Tr}(CQ), \quad (7.15)$$

where C is a symmetric nonnegative definite $n \times n$ matrix, is a generalization of the previous case.

4. The following criterion

$$J = L(Q) = (Qv, v), \quad (7.16)$$

where v is a given nonzero n -vector, is a particular case of (7.15) with

$$C = v * v, \quad C_{ij} = v_i v_j, \quad i, j = 1, \dots, n. \quad (7.17)$$

Here, the symbol $*$ denotes the dyadic product of vectors.

Criterion (7.16) has a clear geometric interpretation: it is related to the projection $\Pi_v(E)$ of the ellipsoid onto the direction of the vector v as follows:

$$\Pi_v(E) = 2(Qv, v)^{1/2}/|v|. \quad (7.18)$$

To prove (7.18), note that the projection $\Pi_v(E)$ is related to the support function $H_E(v)$ of the ellipsoid as follows:

$$\Pi_v(E) = [H_E(v) + H_E(-v)]/|v|. \quad (7.19)$$

Since the support function of the ellipsoid is given by

$$H_E(v) = (a, v) + (Qv, v)^{1/2}, \quad (7.20)$$

equation (7.18) follows immediately from (7.19) and (7.20).

By virtue of (7.18), the minimization of criterion (7.16) is equivalent to the minimization of the projection of the ellipsoid onto the direction of vector v . Other examples of optimality criteria are given in [3].

Below, we consider locally and globally optimal ellipsoids [3, 26].

A smooth family of ellipsoids $E^*(a(t), Q(t))$ is called *locally optimal*, if it is superreachable/subreachable and

$$dL(Q(\tau))/d\tau|_{\tau=t} \rightarrow \min/\max,$$

where the minimum/maximum is taken over all smooth families of superreachable/subreachable ellipsoids $E^\pm(t)$ such that $E^\pm(t) = E^*(t)$.

A smooth family of superreachable/subreachable ellipsoids is called *globally optimal* for a given $t = T$, if the minimum/maximum of $L(Q(T))$ over all superreachable/subreachable ellipsoids is attained on this family.

All definitions and results related to the optimal ellipsoids are true also for the case, where the criterion depends also on time t , so that $J = L(Q, t)$. For example, matrix C in (7.15) and vector v in (7.16) can depend on t : $C = C(t)$, $v = v(t)$.

Note that the volume of an ellipsoid as an optimality criterion (7.14) has a property that singles it out among all other criteria. The optimality of a given ellipsoid in the sense of volume remains intact under any affine transformation in R^n . Thus, the optimality in the sense of volume seems more "basic" property of an approximating ellipsoid than its optimality in the sense of other criteria.

7.5 Equations of Ellipsoids

We consider linear system (7.9) where $u(t)$ is subjected to constraint (7.10) and initial conditions are specified by (7.11).

The center $a^+(t)$ and the matrix $Q^+(t)$ of the outer locally optimal ellipsoid satisfy the following differential equations and initial conditions [3, 5]:

$$\dot{a}^+ = A(t)a^+ + f(t), \quad a^+(s) = a_0, \quad (7.21)$$

$$\dot{Q}^+ = A(t)Q^+ + Q^+A^T(t) + hQ^+ + h^{-1}K, \quad Q^+(s) = Q_0. \quad (7.22)$$

Here, T denotes the transposed matrix, $K(t)$ is expressed via given matrices by the formula

$$K(t) = B(t)G(t)B^T(t), \quad (7.23)$$

and the following notation is used

$$h = \left[\text{Tr} \left(\frac{\partial L}{\partial Q^+} K \right) / \text{Tr} \left(\frac{\partial L}{\partial Q^+} Q^+ \right) \right]^{1/2}. \quad (7.24)$$

Here, Tr is the trace of a matrix, and $\partial L / \partial Q^+$ is a symmetric matrix of partial derivatives $\partial L(Q) / \partial Q_{ij}^+$, $i, j = 1, \dots, n$.

Note that (7.21) for the vector $a(t)$ is linear and does not depend on the chosen optimality criterion $L(Q)$. By contrast, (7.22) for the matrix $Q^+(t)$ is nonlinear and depends on $L(Q)$ via (7.24) for h . For ellipsoids optimal in the sense of volume [see (7.14)], expression (7.24) becomes

$$h = \{n^{-1} \text{Tr} [(Q^+)^{-1} K]\}^{1/2}. \quad (7.25)$$

For the linear criterion (7.15), we have

$$h = [\text{Tr}(CK) / \text{Tr}(CQ^+)]^{1/2}. \quad (7.26)$$

Further simplifications are possible for the criterion (7.16). Substituting C from (7.17) into (7.26), we obtain

$$h = [(Kv, v) / (Q^+v, v)]^{1/2}. \quad (7.27)$$

Equations for inner approximating ellipsoids locally optimal in the sense of volume [3, 5] are as follows:

$$\begin{aligned} \dot{a}^- &= A(t)a^- + f(t), \quad a^-(s) = a_0, \\ \dot{Q}^- &= A(t)Q^- + Q^-A^T(t) + 2K^{1/2}(K^{-1/2}Q^-K^{-1/2})^{1/2}K^{1/2}, \\ Q^-(s) &= Q_0. \end{aligned} \quad (7.28)$$

Here, matrix $K(t)$ is again defined by (7.23). Note that equations for the centers of the inner and outer approximating ellipsoids (7.21) and (7.28) coincide, hence, $a^-(t) \equiv a^+(t)$ for $t \geq s$.

It occurs that (7.28) are true for inner approximating ellipsoids optimal in the sense of all criteria $L(Q)$ satisfying the conditions imposed in the beginning of Sect. 7.4. Thus, these equations have a universal nature.

Let us consider now globally optimal outer ellipsoids. The centers of these ellipsoids coincide with those of locally optimal ones and satisfy the initial value problem (7.21). The matrix $Q^+(t)$ of globally optimal ellipsoids satisfies equation and initial condition (7.22), where matrix K is defined by (7.23), whereas the scalar h is, instead of (7.24), given by the expression

$$h = [\text{Tr}(PK)/\text{Tr}(PQ^+)]^{1/2}. \quad (7.29)$$

Here, $P(t)$ is a symmetric positive definite matrix satisfying the following linear differential equation

$$\dot{P} = -PA(t) - A^T(t)P \quad (7.30)$$

and initial condition at $t = T$:

$$P(T) = [\partial L(Q^+)/\partial Q^+]_{t=T}. \quad (7.31)$$

Hence, we have a two-point boundary value problem for the pair of matrices Q^+ and P described by (7.22), (7.23), (7.29)–(7.31). For the linear criterion (7.15), the initial condition (7.31) is reduced to

$$P(T) = C(T). \quad (7.32)$$

Therefore, for outer ellipsoids globally optimal in the sense of criterion (7.15), the boundary value problem for matrices Q^+ and P becomes decoupled and reduces to two initial value problems: a linear one for $P(t)$ defined by (7.30) and (7.32), and a nonlinear one for $Q^+(t)$ defined by (7.22) and (7.29). First, the problem for $P(t)$ should be solved (from $t = T$ to $t = s$), and then the problem for $Q^+(t)$ (from $t = s$ to $t = T$).

Further simplifications are possible for criterion (7.16). For globally optimal outer ellipsoids, we have, on the strength of (7.17) and (7.32):

$$P(T) = v * v,$$

where v is a given constant n -vector. Let us introduce the adjoint n -vector $\psi(t)$ satisfying the initial value problem:

$$\dot{\psi} = -A^T(t)\psi, \quad \psi(T) = v. \quad (7.33)$$

Then we have [6, 7]

$$P(t) = \psi(t) * \psi(t).$$

Thus, in order to find the matrix $Q^+(t)$ of a globally optimal ellipsoid in the case of criterion (7.16), one is to solve first the linear n -dimensional initial value problem (7.33) for $\psi(t)$ (instead of $n(n+1)/2$ -dimensional problem for $P(t)$ defined by (7.30) and (7.32)), then substitute

$$h = [(K\psi, \psi)/(Q\psi, \psi)]^{1/2}$$

into (7.22) and solve the resultant initial value problem for Q^+ .

Thus, linear optimality criterion (7.15) and, especially, its particular case (7.16) lead to a considerable simplification of equations for optimal outer ellipsoids.

7.6 Transformation of the Equations

In what follows, we restrict ourselves mostly with outer approximating ellipsoids and omit the superscript $+$, so that we denote below: $a^+(t) = a(t)$, $Q^+(t) = Q(t)$.

Equation (7.22) for locally optimal ellipsoids depends on two given matrices: $A(t)$ and $K(t)$, where the symmetric matrix $K(t)$ is given by (7.23). By the change of variables

$$Q = VQ_*V^T, \quad (7.34)$$

where $V(t)$ is an invertible $n \times n$ matrix to be specified below, and Q_* is a new matrix variable, it is possible to simplify (7.22) so that it will contain only one given matrix instead of two.

1. Let us define $\Phi(t)$ as the fundamental matrix of system (7.7):

$$\dot{\Phi} = A(t)\Phi, \quad \Phi(s) = I.$$

Here, I is the $n \times n$ identity matrix.

If we set

$$V(t) = \Phi(t), \quad (7.35)$$

then transformation (7.34) reduces (7.22) to the form

$$\begin{aligned} \dot{Q}_* &= h_* Q_* + h_*^{-1} K_*(t), & Q_*(s) &= Q_0, \\ K_*(t) &= \Phi^{-1}(t) K(t) [\Phi^{-1}(t)]^T. \end{aligned} \quad (7.36)$$

Here, h_* is given by (7.24) where Q should be replaced by $\Phi Q_* \Phi^T$. Explicit expressions [3, 5, 7] for h_* are presented below for criteria (7.14)–(7.16), respectively:

$$\begin{aligned} h_* &= [n^{-1} \text{Tr}(Q_*^{-1} K_*)]^{1/2}, \\ h_* &= [\text{Tr}(C_* K_*) / \text{Tr}(C_* Q_*)]^{1/2}, \quad C_* = \Phi^T C \Phi, \\ h_* &= [(K_* v_*, v_*) / (Q_* v_*, v_*)]^{1/2}, \quad v_* = \Phi^T v. \end{aligned} \quad (7.37)$$

2. Let matrix $K(t)$ from (7.23) be positive definite. By taking

$$V(t) = [K(t)]^{1/2} \quad (7.38)$$

in (7.34), we convert (7.22) to the form

$$\begin{aligned} \dot{Q}_* &= A_* Q_* + Q_* A_*^T + h_* Q_* + h_*^{-1} I, \\ Q_*(s) &= K^{-1/2}(s) Q_0 K^{-1/2}(s). \end{aligned} \quad (7.39)$$

Here, matrix $A_*(t)$ is given by

$$A_*(t) = K^{-1/2}(AK^{1/2} - dK^{1/2}/dt), \quad (7.40)$$

and explicit formulas [7] for h_* are given below for criteria (7.14)–(7.16), respectively:

$$\begin{aligned} h_* &= (n^{-1} \text{Tr} Q_*^{-1})^{1/2}, \\ h_* &= [\text{Tr} C_* / \text{Tr}(C_* Q_*)]^{1/2}, \quad C_* = K^{1/2} C K^{1/2}, \\ h_* &= [(v_*, v_*) / \text{Tr}(Q_* v_*, v_*)]^{1/2}, \quad v_* = K^{1/2} v. \end{aligned}$$

Thus, by choosing matrix V in (7.34) according to (7.35), we reduce (7.22) for matrix Q to the form (7.36) that corresponds to the case where $A = 0$. On the other hand, by taking matrix V in accordance with (7.38), we come to (7.39) where K is replaced by the unity matrix I . Therefore, we can restrict ourselves with considering (7.22) only in the cases where either $A = 0$ or $K = I$.

Similar simplifications take place also for (7.28) for inner ellipsoids.

Let us consider now equations for globally optimal outer ellipsoids [7] and use the change of variables (7.34) for Q and

$$P = (V^{-1})^T P_* V^{-1} \quad (7.41)$$

for P , where P_* is a new variable. Let us restrict ourselves with the case of linear criteria (7.15) and (7.16). If V is defined by (7.35), we obtain from (7.30) and (7.32) for criterion (7.15):

$$P_*(t) = \Phi^T(T) C(T) \Phi(T) = \text{const.}$$

As a result, the equation for matrix $Q_*(t)$ coincides with (7.36), where

$$h_* = [\text{Tr}(P_*K_*)/\text{Tr}(P_*Q_*)]^{1/2}.$$

For criterion (7.16), h_* is determined by (7.37) with $v_* = \Phi^T(T)v$.

By defining V in (7.34) and (7.41) in accordance with (7.38), we again obtain for criterion (7.15), equations (7.39) for Q_* , equations

$$\dot{P}_* = -P_*A_* - A_*^T P_*, \quad P_*(T) = K^{1/2}(T)C(T)K^{1/2}(T)$$

for P_* , and the same expression (7.40) for A_* .

For criterion (7.16), (7.39) for Q_* and (7.40) for A_* still hold. Here, h_* is defined by

$$h_* = [(\psi_*, \psi_*)/(Q_*\psi_*, \psi_*)]^{1/2},$$

where the adjoint vector $\psi_*(t)$ satisfies the following initial value problem that replaces (7.33):

$$\dot{\psi}_* = -A_*^T \psi_*, \quad \psi_*(T) = K^{1/2}(T)v.$$

7.7 Properties of Optimal Ellipsoids

Outer approximating ellipsoids $E(a(t), Q(t))$ optimal in the sense of criterion (7.16) have the following properties:

1. Globally optimal ellipsoids touch reachable sets $D(t, s, M)$ for all $t \in [s, T]$ at points $x(t)$, where the normal to the boundary of these sets is parallel to the vector $\psi(t)$ defined by (7.33) [6]. In other words, these ellipsoids are tight in the sense of [16].
2. Globally optimal ellipsoids are also locally optimal for the vector $v(t) = \psi(t)$.
3. Locally optimal ellipsoids for the vector $v(t)$ defined by the initial value problem:

$$v(t) = \psi(t), \quad \dot{\psi} = -A^T(t)\psi, \quad \psi(s) = v^0, \quad (7.42)$$

where v^0 is an arbitrary vector, are also globally optimal for any terminal time instant $T \geq s$ and for the criterion $J = (Qv(T), v(T))$.

To construct these locally (and also globally) optimal ellipsoids, one is to solve the linear initial value problem (7.21) for $a^+(t)$ and also initial value problem consisting of (7.22) for Q^+ and (7.42) for $\psi(t)$. Here, the initial vector v^0 can be chosen arbitrarily, and different vectors v^0 correspond to different approximating ellipsoids touching reachable sets at different points.

Various properties of nonlinear equations (7.22) and (7.28) governing the evolution of locally optimal ellipsoids have been studied [1–3, 5–7, 26–28].

As a rule, the nonlinear differential equations for ellipsoids are to be integrated numerically. However, a number of explicit analytical solutions have been obtained both for locally [1–3, 5] and globally [6, 26–28] optimal ellipsoids.

Asymptotic behavior of the solution of (7.22) and (7.28) have been analyzed in the vicinity of the initial point $t = s$, if $Q_0 = 0$ [2, 3]. This case corresponds to the situation, where the initial set M in (7.11) is a given point $x(s) = a_0$. In this important case, (7.22) and (7.28) have a singularity [see also (7.24)–(7.27)], and the obtained asymptotic expansions of the solution are needed to start the numerical integration of equations near the initial point $t = s$ for the case where $Q_0 = 0$.

Also, asymptotic behavior for solutions of (7.22) and (7.28) at infinity ($t \rightarrow \infty$) have been analyzed [2, 3, 6, 26–28].

7.8 Generalizations

The method of ellipsoids has been extended to the case, where the parameters of the linear system (7.9) are uncertain and/or subjected to unknown but bounded perturbations. Consider the following system

$$\dot{x} = [A_0(t) + A_1(t)]x + f(t), \quad (7.43)$$

where $x \in R^n$ is the state, matrix $A_0(t)$ and n -vector $f(t)$ are given functions of time, whereas the matrix $A_1(t)$ is unknown, and its elements $a_{ij}^1(t)$ are bounded:

$$|a_{ij}^1(t)| \leq b_{ij}, \quad i, j = 1, \dots, n, \quad t \geq s. \quad (7.44)$$

Here, b_{ij} are given nonnegative numbers. The system described by (7.43) and (7.44) models the situation, where some parameters of the system are uncertain (fixed but unknown) or changeable, for example, in the case of parametric excitation.

Outer ellipsoidal estimates $E(a(t), Q(t))$ on the reachable sets of the system described by (7.43) and (7.44) have been obtained [4].

Suppose the initial data are given by (7.11). The equation for the center $a(t)$ of the approximating ellipsoid is still the same as (7.21), where $A(t)$ is replaced by $A_0(t)$, so that we have

$$\dot{a} = A_0(t)a + f(t), \quad a(s) = a_0.$$

Nonlinear matrix equation for $Q(t)$ differs from (7.22) and has the form

$$\begin{aligned} \dot{Q} &= A_0 Q + Q A_0^T + h Q + h^{-1} R(a, Q), \\ h &= [n^{-1} \text{Tr}(Q^{-1} R)]^{1/2}, \quad R = \text{diag}(R_1^2, \dots, R_n^2), \\ R_i &= \sum_{j=1}^n b_{ij} |a_j| + \left(\max_{\sigma} \sum Q_{jk} b_{ij} b_{ik} \sigma_{ij} \sigma_{ik} \right)^{1/2}. \end{aligned} \quad (7.45)$$

Here, the maximum is taken over all $\sigma_{ij} = \pm 1$, $i, j = 1, \dots, n$. Note that, in contrast to (7.22), the right-hand side of (7.45) for Q depends on vector a .

The method of ellipsoids can be extended also to nonlinear systems [2, 3]. The main idea is to construct a linear comparison system described by (7.9), (7.10), and (7.11), so that all possible motions of the original nonlinear system are within the reachable sets of the linear one. For example, consider a nonlinear system

$$\dot{x} = A(t)x + \varphi(u, t), \quad |\varphi(u, t)| \leq \varphi_0(t), \quad t \geq s, \quad (7.46)$$

where u is the disturbance, and the absolute value of the nonlinearity $\varphi(u, t)$ is bounded by $\varphi_0(t)$ for all admissible u and $t \geq s$: $|\varphi(u, t)| \leq \varphi_0(t)$.

Then, the following linear system

$$\dot{x} = A(t)x + w, \quad w \in E(0, \varphi_0^2(t)I) \quad (7.47)$$

can serve as a comparison system for (7.46). Here, w is a disturbance bounded by a ball.

An outer approximating ellipsoid $E(a(t), Q(t))$ for linear system (7.47) will provide an outer bound also for reachable sets of the original nonlinear system (7.46).

Similarly, inner ellipsoidal bounds for reachable sets of nonlinear systems can be obtained [2, 3].

The class of approximating sets can be extended: besides ellipsoids, also intersections and unions of several ellipsoids can be considered [2, 3]. Thus, the approximation of reachable sets can be improved significantly.

7.9 Applications

Two-sided ellipsoidal approximations of reachable sets can be used for the solution and approximation of various problems in control and state estimation. Here, we will only briefly mention these applications; see [3, 5] for details.

7.9.1 Two-Sided Estimates in Optimal Control

Consider the following optimal control problem for system (7.9) under the initial condition $x(s) = x^0$.

Find the control subject to constraint (7.10) that provides the minimum of the given terminal functional $J = F(x(T))$ at $t = T > s$.

Let us obtain two-sided approximating ellipsoids (locally or globally optimal) and evaluate the following minima of the function $F(x)$ over the ellipsoids

$$F^\pm = \min F(x), \quad x \in E(a^\pm(T), Q^\pm(T)).$$

Then the two-sided bounds $F^+ \leq \min J \leq F^-$ are true for the required minimum of the functional J . To obtain these bounds, one needs, first, to find the inner and outer ellipsoids, and, second, to solve the problems of nonlinear programming, namely, to minimize the function $F(x)$ over the ellipsoidal sets. Note that the first part of this procedure does not depend on the function $F(x)$; if we change this function, we are to change only the second part of the procedure.

7.9.2 Two-Sided Bounds on Time for the Time-Optimal Problem

Let the time-optimal problem ($T \rightarrow \min$) is set up for the system (7.9) under the initial condition $x(s) = x_0$. The terminal state is fixed: $x(T) = x^*$. Let us find the minimal time instants T^+ and T^- when the ellipsoids $E(a^\pm(t), Q^\pm(t))$ contain the point x^* :

$$T^\pm = \min t, \quad x^* \in E(a^\pm(t), Q^\pm(t)), \quad t \geq s.$$

Then the two-sided bounds $T^+ \leq T \leq T^-$ for T are true.

7.9.3 Suboptimal Control

Consider again a linear system described by (7.9) and (7.10) under the initial condition $x(s) = x^0$. Suppose that the given terminal state $x(T) = x^*$ at some instant T belongs to the inner ellipsoid $E(a^-(T), Q^-(T))$. Then there exists an admissible control $u(t)$ bringing the system to this terminal state at $t = T$. This control can be constructed efficiently [12, 13] and is defined by the following procedure. First, find the solution of the initial value problems (7.28) for functions $a^-(t)$ and $Q^-(t)$ that determine the inner approximating ellipsoid.

Denote

$$R(t) = K^{1/2}(K^{-1/2}Q^-K^{-1/2})^{1/2}K^{1/2}$$

and solve the auxiliary initial value problem

$$\begin{aligned} \dot{\psi} &= -A^T \psi - (Q^-)^{-1} R \psi \quad t \in [s, T], \\ \psi(T) &= [Q^-(T)]^{-1} [x^* - a^-(T)]. \end{aligned}$$

Then the admissible control $u(t)$ bringing system (7.9) from the initial state $x(s) = x^0$ to the prescribed terminal state $x(T) = x^*$ is given by the expression

$$u(t) = R(t)\psi(t), \quad t \in [s, T].$$

This control can be called suboptimal since, if the terminal time T is equal to the upper bound T^- for the time-optimal problem introduced in Sect. 7.9.2, this control brings our system to the prescribed terminal state at $t = T = T^-$.

7.9.4 Differential Games

Consider now a differential game of two players X and Y described by equations similar to (7.9) and (7.10):

$$\begin{aligned} X : \quad & \dot{x} = A_x(t)x + B_x(t)u + f_x(t), \\ & u \in E(0, G_x(t)), \quad x(s) = x^0; \\ Y : \quad & \dot{y} = A_y(t)y + B_y(t)v + f_y(t), \\ & v \in E(0, G_y(t)), \quad y(s) = y^0. \end{aligned}$$

Here, x and y are the state vectors of the players X and Y , respectively, u and v are their controls, and x^0 and y^0 are the respective initial states. The cost functional J is a given scalar function of the terminal states of the players at the prescribed time instant T :

$$J = \Phi(x(T), y(T)), \quad T > s.$$

Player X seeks to minimize J while player Y , which can be also identified with uncertain disturbances, opposes Y and tends to maximize J . The following inequalities

$$\max_{y \in D_y} \min_{x \in D_x} \Phi(x, y) \leq J^* \leq \min_{x \in D_x} \max_{y \in D_y} \Phi(x, y),$$

where D_x and D_y are the reachable sets of players X and Y , respectively, at the instant T , provide evident two-sided bounds on the optimal value J^* of the functional J that corresponds to optimal strategies of both players.

Using ellipsoidal bounds

$$E_x^\pm = E(a_x^\pm(t), Q_x^\pm(t)), \quad E_y^\pm = E(a_y^\pm(t), Q_y^\pm(t))$$

on reachable sets for players X and Y , we obtain the following two-sided estimates on J^* :

$$\Phi_1 \leq J^* \leq \Phi_2, \quad \Phi_1 = \max_{y \in E_y^-} \min_{x \in E_x^+} \Phi(x, y), \quad \Phi_2 = \min_{x \in E_x^-} \max_{y \in E_y^+} \Phi(x, y). \quad (7.48)$$

Suppose the pairs of points x_1^*, y_1^* and x_2^*, y_2^* are found where the maximin Φ_1 and minimax Φ_2 from (7.48) are attained, respectively. Using results of Sect. 7.9.3,

we can find the control $u(t)$ bringing player X to the state $x_2^* \in E_x^-$ at $t = T$, and also the control $v(t)$ bringing player Y to the state $y_1^* \in E_y^-$ at $t = T$. If player X applies the open-loop control $u(t)$, then the value of functional J does not exceed Φ_2 under any admissible control of player Y . On the other hand, if player Y applies the open-loop control $v(t)$, then the value of functional J is not less than Φ_1 under any admissible control of player X . Thus, using approximating ellipsoids, we can obtain two-sided bounds on the value of the cost functional and determine open-loop controls of the players that ensure these bounds.

Approximating ellipsoids can be also used for obtaining two-sided bounds in games of pursuit-evasion. In this context, the rule of extremal aiming [14, 15] is used. Numerical example is presented in [11].

7.9.5 Control of Uncertain Systems

Consider a system subjected to both the control $u(t)$ and disturbance $v(t)$:

$$\dot{x} = A(t)x + B(t)u + C(t)v + f(t), \quad t \geq a. \quad (7.49)$$

Knowing the bound $v(t) \in E(0, G(t))$ on the disturbance, we can obtain the outer bound $x(t) \in E(a(t), Q(t))$ on all trajectories of system (7.49) subjected to the given control $u(t)$. Then, equations (7.21) for $a(t)$ and (7.22) for $Q(t)$ become

$$\dot{a} = Aa + Bu + f, \quad \dot{Q} = AQ + QA^T + hQ + h^{-1}K, \quad K = BGB^T$$

and can be considered as a control system for the whole set of possible trajectories. Various control problems can be set up for this system, and different control methods can be applied to these problems.

7.9.6 Other Applications

In case where $u(t)$ is a bounded disturbance in (7.9), the outer approximating ellipsoid provides an estimate on possible deviations of the trajectory caused by the disturbance. Such estimates are sometimes associated with the notion of “practical stability”. For example, possible deviation of the trajectory of a moving body in the presence of wind disturbances can be evaluated.

Various applications of ellipsoidal bounds to parameter estimation are considered in [30, 31]. Aerospace applications of approximating ellipsoids are discussed in [25, 32].

7.9.7 State Estimation in the Presence of Observation Errors

Consider again the system subjected to uncertain disturbances and described by (7.9), (7.10), and (7.11). Suppose that, at the given time instants t_i , the results of observations

$$y_i = H_i x(t_i) + \xi_i, \quad t_i \geq s, \quad i = 0, 1, \dots, \tag{7.50}$$

become available. Here, y_i are m -vectors of observation results, H_i are given $m \times n$ matrices, and ξ_i are m -dimensional observation errors subject to constraints $\xi_i \in E(0, L_i)$, where L_i are given symmetric positive definite $m \times m$ matrices. Thus, at the instants t_i , the state $x(t_i)$ of the system belongs to the intersection of two ellipsoidal sets:

$$\begin{aligned} x(t) &\in E(t_i) \cap \tilde{E}_i, \\ \tilde{E}_i &= \{x : (L_i^{-1}[H_i x(t_i) - y_i], [H_i x(t_i) - y_i]) \leq 1\}. \end{aligned} \tag{7.51}$$

Here, $E(t_i)$ is the outer ellipsoidal state estimate of the system based on all information available for $t < t_i$, and \tilde{E}_i is the ellipsoid corresponding to the observation (7.50) at $t = t_i$.

To design the process of the ellipsoidal state estimation, we are to construct the ellipsoid $E(t_i+0)$ that contains the intersection of ellipsoids (7.51). Then we can use $E(t_i+0)$ as the initial ellipsoid for the next time interval (t_i, t_{i+1}) , see Fig. 7.2. It is desirable to find $E(t_i+0)$ that minimizes certain optimality criterion (see Sect. 7.4). Using optimal and suboptimal outer ellipsoidal bounds for the intersection of two ellipsoids [2, 3], the recursive procedure for the ellipsoidal state estimation in the presence of observation errors has been developed. The state estimation procedure

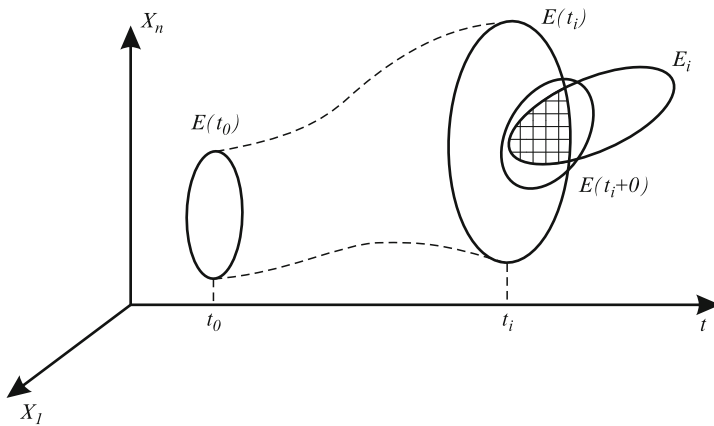


Fig. 7.2 Ellipsoidal state estimation

has been extended also to the case of continuous observations. Thus, the guaranteed analogue of the well-known Kalman filtering has been elaborated [2, 3, 5].

Applications of ellipsoidal technique to state estimation are considered in [17, 18, 25, 30, 31].

7.10 Ellipsoidal vs. Interval Analysis

There exists a vast literature on the interval analysis that is widely used in the computational mathematics [8, 9, 20, 22]. In the interval analysis, to obtain the guaranteed error estimates, one operates with intervals $[x^-, x^+]$, instead of precise value x that is unknown but bounded, $x \in [x^-, x^+]$. When this approach is applied to vectors $x = (x_1, \dots, x_n)$, one is to deal with boxes

$$B = \{x : x_i^- \leq x \leq x_i^+, \quad i = 1, \dots, n\}, \quad (7.52)$$

i.e., rectangular parallelepipeds with sides parallel to coordinate axes.

The interval approach and its generalizations, where uncertainty domains are bounded by various polytopes, are also applied in the control and identification, see [10, 19, 21, 23, 24, 34].

Let us discuss and compare the method of ellipsoids with the interval methods.

One can consider several levels of generalization of interval methods. The sets of uncertainty can be bounded by: (a) boxes (7.52); (b) rectangular parallelepipeds; (c) arbitrary parallelepipeds; (d) polytopes. The class (d) of arbitrary polytopes seems to be too wide. As already mentioned in Sect. 7.2, the class (c), like the class of ellipsoids, is invariant with respect to linear transformations, but it requires almost two times more parameters for its description than the class of ellipsoids. The classes (a) and (b) are not invariant with respect to linear transformations.

In this context, the class of boxes (7.52) seems to be more attractive among classes (a)–(d) because it has the simplest description and requires only $2n$ parameters.

However, this class can lead to an undesirable but essential error increase.

Consider a simplest vectorial operation $y = Ax$, where x is defined by a box (7.52) and A is a given $n \times n$ matrix. As a result of this operation, box (7.52) is transformed to a parallelepiped P . To obtain a box B' for vector y , one needs to take a box that contains P , $B' \supset P$, and thus to expand the set of possible vectors y . In other words, the error estimate will become wider. In [29], the results of operation $y = Ax$ have been analyzed both for ellipsoidal and interval uncertainty bounds. It has been shown that the ellipsoidal bounds are frequently superior to interval ones and lead to tighter error estimates than those given by the interval analysis.

Thus, it seems that the method of ellipsoids can be of use for error estimation in numerical analysis.

7.11 Conclusions

The method of ellipsoids seems to be an efficient technique for the analysis of dynamical systems subjected to uncertain perturbations and observation errors. By means of this approach, exact and approximate solutions as well as reliable two-sided bounds for a number of basic problems in control and estimation can be obtained.

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Chapter 8

Expected Total Cost Minimum Design of Plane Frames by Means of Stochastic Linear Programming Methods

Kurt Marti

Abstract Yield stresses, allowable stresses, moment capacities (plastic moments with respect to compression, tension and rotation), applied loadings, cost factors, manufacturing errors, etc., are not given fixed quantities in structural analysis and optimal design problems, but must be modeled as random variables with a certain joint probability distribution. Problems from plastic analysis and optimal plastic design are based on the convex yield (feasibility) criterion and the linear equilibrium equation for the stress (state) vector.

After the formulation of the basic mechanical conditions including the relevant material strength parameters and load components as well as the involved design variables (as, e.g., sizing variables) for plane frames, several approximations are considered: (1) approximation of the convex yield (feasible) domain by means of convex polyhedrons (piecewise linearization of the yield domain); (2) treatment of symmetric and non symmetric yield stresses with respect to compression and tension; (3) approximation of the yield condition by using given reference capacities.

As a result, for the survival of plane frames a certain system of necessary and/or sufficient linear equalities and inequalities is obtained. Evaluating the recourse costs, i.e., the costs for violations of the survival condition by means of piecewise linear convex cost functions, a linear program is derived for the minimization of the total costs including weight-, volume- or more general initial (construction) costs. Appropriate cost factors are given. Considering then the expected total costs from construction as well as from possible structural damages or failures, a stochastic linear optimization problem (SLP) is obtained. Finally, discretization of the probability distribution of the random parameter vector yields a (large scale) linear program (LP) having a special structure. For LP's of this type numerous, very efficient LP-solvers are available – also for the solution of very large scale problems.

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8.1 Introduction

8.1.1 *Plastic Analysis of Structures*

Many materials, e.g., most of metals, have distinct, plastic properties, i.e., they are ductile, see, e.g., [10,23]. Even after the stress intensity attains the yield point stress, such materials can deform considerably without breaking. This implies that if the stress intensity at a certain point of a hyperstatic structure reaches the critical (yield) value, the structure does not necessarily fail or deform excessively. Instead, a certain amount of stress redistribution takes place and some further load increments can be supported. Structural failure does not occur before a kinematic mechanism of unconstrained plastic flow develops. Thus, the actual load-carrying capacity of a structure is higher (in some cases quite considerably) than that derived from classical elastic analysis. A crucial question for the engineer designing structures like buildings, bridges, etc., or structural components is to which extent a plastic deformation is permissible without leading to a failure of the structure, the component, resp., with respect to the expected load and material strength conditions. Applying standard methods, the load carrying capacity is determined using a certain code with general rules for safety evaluations. The use of such general rules may be very expensive in the safety evaluation and design of a structure. On the other hand, safety assessment and design based on stochastic optimization techniques, taking into account the available knowledge about random parameter variations, reduce the expected total project costs (primary costs, e.g., costs of construction, plus recourse costs, e.g., strengthening costs) considerably. Consequently, this way one obtains more robust (safe) information about the maximum load factors, hence, the carrying capacity, as well as about robust optimal designs. A further big advantage of stochastic optimization methods is the possibility of updates of the maximum load factors and robust optimal designs based on inspection, sampling and other posterior information about the probability distribution of the random parameters. For elastic-perfectly plastic materials, the ultimate load condition corresponding to complete collapse of the structure can be obtained through application of a pair of dual theorems [14, 18, 23]:

(ST) Static Theorem (lower bound or safe theorem) If any stress distribution throughout the structure can be found which is everywhere in equilibrium internally and balances the external loads, and at the same time does not violate the yield condition, those external loads will be carried safely by the structure.

(KT) Kinematic Theorem (upper bound or unsafe theorem)

Collapse occurs if a collapse mechanism, fulfilling the compatibility condition, exists such that the work done by the external loads is larger than the corresponding internal plastic work.

Limit analysis is concerned [5, 8, 10, 14, 16, 18, 19, 22, 24, 33, 34, 36, 37, 40, 41] with establishing the strength of a structure, i.e., its capacity for the supporting of loads. Hence, using the plastic ductility of structural materials in

improving the design of structures, limit analysis is not concerned with deformation: it can not therefore provide the load carrying capacity for a structure with elements that have a limited ductility or deformability, nor for a structure which becomes unstable because of the displacements induced by plastic deformation, see [10, 13, 18, 24].

8.1.2 Limit (Collapse) Load Analysis of Structures as a Linear Programming Problem

Assuming that the material behaves as an elastic-perfectly plastic material [17, 23, 32] a conservative estimate of the collapse load factor λ_T is based [5–8, 10, 13, 14, 22, 33, 40] on the following linear program:

$$\text{maximize } \lambda \tag{8.1a}$$

s.t.

$$F^L \leq F \leq F^U \tag{8.1b}$$

$$CF = \lambda R_0. \tag{8.1c}$$

Here, (8.1c) is the equilibrium equation of a statically indeterminate loaded structure involving an $m \times n$ matrix $C = (c_{ij}), m < n$, of given coefficients $c_{ij}, 1 \leq i \leq m, 1 \leq j \leq n$, depending on the undeformed geometry of the structure having n_0 members (elements). After taking into account the given support (boundary) conditions, we may suppose that $\text{rank } C = m$. Furthermore, R_0 is an external load m -vector, and F denotes the n -vector of internal forces and bending-moments in the relevant points (sections, nodes or elements) of lower and upper bounds F^L, F^U .

For a plane or spatial truss [25, 38] we have that $n = n_0$, the matrix C contains the direction cosines of the members, and F involves only the normal (axial) forces moreover,

$$F_j^L := \sigma_{yj}^L A_j, F_j^U := \sigma_{yj}^U A_j, j = 1, \dots, n(= n_0), \tag{8.2}$$

where A_j is the (given) cross-sectional area, and $\sigma_{yj}^L, \sigma_{yj}^U$, respectively, denotes the yield stress in compression (negative values) and tension (positive values) of the j -th member of the truss. In case of a plane frame, F is composed of subvectors [38],

$$F^{(k)} = \begin{pmatrix} F_1^{(k)} \\ F_2^{(k)} \\ F_3^{(k)} \end{pmatrix} = \begin{pmatrix} t_k \\ m_k^+ \\ m_k^- \end{pmatrix}, \tag{8.3a}$$

where $F_1^{(k)} = t_k$ denotes the normal (axial) force, and $F_2^{(k)} = m_k^+, F_3^{(k)} = m_k^-$ are the bending-moments at the positive (“right”), negative (“left”) end of the k -th member with respect to a certain chosen orientation of the members. In this case

F^L, F^U contain – for each member k – the subvectors

$$F^{(k)L} = \begin{pmatrix} \sigma_{yk}^L A_k \\ -M_{kpl} \\ -M_{kpl} \end{pmatrix}, F^{(k)U} = \begin{pmatrix} \sigma_{yk}^U A_k \\ M_{kpl} \\ M_{kpl} \end{pmatrix}, \quad (8.3b)$$

resp., where $M_{kpl}, k = 1, \dots, n_0$, denotes the plastic moments (moment capacities) [17, 32] given by

$$M_{kpl} = \sigma_{yk}^U W_{kpl}, \quad (8.3c)$$

and $W_{kpl} = W_{kpl}(A_k)$ is the plastic section modulus of the cross-section of the k -th member (beam) with respect to the local z -axis. In order to omit instabilities, such as buckling, σ_{yk}^L can be selected by

$$\sigma_{yk}^L := -\kappa_k \sigma_{yk}^U \quad (8.3d)$$

with a certain reduction factor κ_k .

For a *space frame* [25, 38], corresponding to the k -th member (beam), F contains the subvector

$$F^{(k)} := (t_k, m_{kT}, m_{k\bar{y}}^+, m_{k\bar{z}}^+, m_{k\bar{y}}^-, m_{k\bar{z}}^-)^T, \quad (8.4a)$$

where t_k is the normal (axial) force, m_{kT} the twisting moment, and $m_{k\bar{y}}^+, m_{k\bar{z}}^+, m_{k\bar{y}}^-, m_{k\bar{z}}^-$ denote four bending moments with respect to the local \bar{y}, \bar{z} -axis at the positive, negative end of the beam, respectively. Finally, the bounds F^L, F^U for F are given by

$$F^{(k)L} = (\sigma_{yk}^L A_k, -M_{kpl}^{\bar{p}}, -M_{kpl}^{\bar{y}}, -M_{kpl}^{\bar{z}}, -M_{kpl}^{\bar{y}}, -M_{kpl}^{\bar{z}})^T \quad (8.4b)$$

$$F^{(k)U} = (\sigma_{yk}^U A_k, M_{kpl}^{\bar{p}}, M_{kpl}^{\bar{y}}, M_{kpl}^{\bar{z}}, M_{kpl}^{\bar{y}}, M_{kpl}^{\bar{z}})^T, \quad (8.4c)$$

where [17, 32]

$$M_{kpl}^{\bar{p}} := \tau_{yk} W_{kpl}^{\bar{p}}, M_{kpl}^{\bar{y}} := \sigma_{yk}^U W_{kpl}^{\bar{y}}, M_{kpl}^{\bar{z}} := \sigma_{yk}^U W_{kpl}^{\bar{z}}, \quad (8.4d)$$

are the plastic moments of the cross-section of the k -th element with respect to the local twisting axis, the local \bar{y}, \bar{z} -axis, respectively. In (8.4d), $W_{kpl}^{\bar{p}} = W_{kpl}^{\bar{p}}(x)$ and $W_{kpl}^{\bar{y}} = W_{kpl}^{\bar{y}}(x), W_{kpl}^{\bar{z}} = W_{kpl}^{\bar{z}}(x)$, resp., denote the polar, axial modulus of the cross-sectional area of the k -th beam and τ_{yk} denotes the yield stress with respect to torsion; we suppose that $\tau_{yk} = \kappa_{\tau k} \sigma_{yk}^U$ with a reduction factor $\kappa_{\tau k}$. Moreover, x denotes the r -vector of design variables, see Sect. 8.1.1 for more details.

Remark 8.1. Possible plastic hinges [17, 24, 32] are taken into account by inserting appropriate eccentricities $e_{kl} > 0, e_{kr} > 0, k = 1, \dots, n_0$, with $e_{kl}, e_{kr} \ll L_k$, where L_k is the length of the k -th beam.

Remark 8.2. Working with more general yield polygons [1, 40, 42], the stress condition (8.1b) is replaced by the more general system of inequalities

$$H(F_d^U)^{-1} F \leq h. \quad (8.5a)$$

Here, (H, h) is a given $v \times (n + 1)$ matrix, and $F_d^U := (F_j^U \delta_{ij})$ denotes the $n \times n$ diagonal matrix of principal axial and bending plastic capacities

$$F_j^U := \sigma_{ykj}^U A_{kj}, F_j^U := \sigma_{ykj}^U W_{kjp}^{\kappa j}, \quad (8.5b)$$

where $kj, \kappa j$ are indices as arising in (8.3b)–(8.4d). The more general case (8.5a) can be treated by similar methods as the case (8.1b) which is considered here.

8.1.3 Plastic and Elastic Design of Structures

In the plastic design of trusses and frames [22, 26, 27, 29, 34, 36] having n_0 members, the n -vectors F^L, F^U of lower and upper bounds

$$F^L = F^L(\sigma_y^L, \sigma_y^U, x), F^U = F^U(\sigma_y^L, \sigma_y^U, x), \quad (8.6)$$

for the n -vector F of internal member forces and bending moments $F_j, j = 1, \dots, n$, are determined [13, 22] by the yield stresses, i.e., compressive limiting stresses (negative values) $\sigma_y^L = (\sigma_{y1}^L, \dots, \sigma_{yn_0}^L)^T$, the tensile yield stresses $\sigma_y^U = (\sigma_{y1}^U, \dots, \sigma_{yn_0}^U)^T$, and the r -vector

$$x = (x_1, x_2, \dots, x_r)^T \quad (8.7)$$

of design variables of the structure. In case of trusses we have that, cf. (8.2),

$$\begin{aligned} F^L &= \sigma_{yd}^L A(x) = A(x)_d \sigma_y^L, \\ F^U &= \sigma_{yd}^U A(x) = A(x)_d \sigma_y^U, \end{aligned} \quad (8.8)$$

where $n = n_0$, and $\sigma_{yd}^L, \sigma_{yd}^U$ denote the $n \times n$ diagonal matrices having the diagonal elements $\sigma_{yj}^L, \sigma_{yj}^U$, respectively, $j = 1, \dots, n$, moreover,

$$A(x) = [A_1(x), \dots, A_n(x)]^T \quad (8.9)$$

is the n -vector of cross-sectional area $A_j = A_j(x), j = 1, \dots, n$, depending on the r -vector x of design variables $x_\kappa, \kappa = 1, \dots, r$, and $A(x)_d$ denotes the $n \times n$ diagonal matrix having the diagonal elements $A_j = A_j(x), 1 < j < n$.

Corresponding to (8.1c), here the equilibrium equation reads

$$CF = R_u, \quad (8.10)$$

where R_u describes [22] the ultimate load [representing constant external loads or self-weight expressed in linear terms of $A(x)$].

The *plastic design* of structures can be represented then [1,2] by the optimization problem

$$\min G(x), \quad (8.11a)$$

s.t.

$$F^L(\sigma_y^L, \sigma_y^U, x) \leq F \leq F^U(\sigma_y^L, \sigma_y^U, x) \quad (8.11b)$$

$$CF = R_u \quad (8.11c)$$

$$x \in D, \quad (8.11d)$$

where $G = G(x)$ is a certain objective function, e.g., the volume or weight of the structure, and $C \subset \mathbb{R}^+$ denotes the convex set of admissible design vectors x .

Remark 8.3. As mentioned in Remark 8.2, working with more general yield polygons, (8.11b) is replaced by the condition

$$H[F^U(\sigma_y^U, x)_d]^{-1} F \leq h. \quad (8.11e)$$

For the *elastic design* we must replace the yield stresses σ_y^L, σ_y^U by the allowable stresses σ_a^L, σ_a^U and instead of ultimate loads we consider service loads R_s . Hence, instead of (8.11a–d) we have the related program

$$\min G(x), \quad (8.12a)$$

s.t.

$$F^L(\sigma_a^L, \sigma_a^U, x) \leq F \leq F^U(\sigma_a^L, \sigma_a^U, x), \quad (8.12b)$$

$$CF = R_s, \quad (8.12c)$$

$$x^L \leq x \leq x^U, \quad (8.12d)$$

where x^L, x^U still denote lower and upper bounds for x .

8.2 Plane Frames

For each bar $i = 1, \dots, B$ of a plane frame with member load vector $F_i = (t_i, m_i^+, m_i^-)^T$ we consider [37, 41] the load at the negative, positive end

$$F_i^- := (t_i, m_i^-)^T, F_i^+ := (t_i, m_i^+)^T, \quad (8.13)$$

respectively.

Furthermore, for each bar/beam with rigid joints we have several plastic capacities: The plastic capacity N_{ipl}^L of the bar with respect to axial compression, hence, the maximum axial force under compression is given by

$$N_{ipl}^L = |\sigma_{yi}^L| \cdot A_i, \quad (8.14a)$$

where $\sigma_{yi}^L < 0$ denotes the (negative) yield stress with respect to compression and A_i is the cross sectional area of the i th element. Correspondingly, the plastic capacity with respect to (axial) tension reads:

$$N_{ipl}^U = \sigma_{yi}^U \cdot A_i, \quad (8.14b)$$

where $\sigma_{yi}^U > 0$ is the yield stress with respect to tension. Besides the plastic capacities with respect to the normal force, we have the moment capacity

$$M_{ipl} = \sigma_{yi}^U \cdot W_{ipl} \quad (8.14c)$$

with respect to the bending moments at the ends of the bar i .

Remark 8.4. Note that all plastic capacities have nonnegative values.

Using the plastic capacities (8.14a–c), the load vectors F_i^+ , F_i^- given by (8.13) can be replaced by dimensionless quantities

$$F_i^{L-} := \left(\frac{t_i}{N_{ipl}^L}, \frac{m_i^-}{M_{ipl}} \right)^T, \quad F_i^{U-} := \left(\frac{t_i}{N_{ipl}^U}, \frac{m_i^-}{M_{ipl}} \right)^T \quad (8.15a,b)$$

$$F_i^{L+} := \left(\frac{t_i}{N_{ipl}^L}, \frac{m_i^+}{M_{ipl}} \right)^T, \quad F_i^{U+} := \left(\frac{t_i}{N_{ipl}^U}, \frac{m_i^+}{M_{ipl}} \right)^T \quad (8.15c,d)$$

for the negative, positive end, resp., of the i th bar.

Remark 8.5 (Symmetric yield stresses under compression and tension).

In the important special case that the absolute values of the yield stresses under compression (<0) and tension (>0) are equal, hence,

$$\sigma_{yi}^L = -\sigma_{yi}^U \quad (8.16a)$$

$$N_{ipl}^L = N_{ipl}^U =: N_{ipl}. \quad (8.16b)$$

The limit between the elastic and plastic state of the elements is described by the feasibility or yield condition: At the negative end we have the condition

$$F_i^{L-} \in K_i, \quad F_i^{U-} \in K_i \quad (8.17a,b)$$

and at the positive end the condition reads

$$F_i^{L+} \in K_i, F_i^{U+} \in K_i. \tag{8.17c,d}$$

Here, $K_i, K_i \subset \mathbb{R}^2$, denotes the feasible domain of bar “i” having the following properties:

- K_i is a closed, convex subset of \mathbb{R}^2 .
- The origin 0 of \mathbb{R}^2 is an interior point of K_i .
- The interior $\overset{\circ}{K}_i$ of K_i represents the elastic states.
- At the boundary ∂K_i yielding of the material starts.

Considering, e.g., bars with rectangular cross sectional areas and symmetric yield stresses, cf. Remark 8.5, K_i is given by $K_i = K_{0,sym}$, where [19,21]

$$K_{0,sym} = \{(x, y)^T : x^2 + |y| \leq 1\} \tag{8.18}$$

where $x = \frac{N}{N_{pl}}$ and $y = \frac{M}{M_{pl}}$, (see Fig. 8.1). Note that the symbols $x, y, (x, y)$, resp., denote in this Sect. 8.2 just real variables, a point in the real plane.

In case (8.18) and supposing symmetric yield stresses, the yield condition (8.17a–d) reads

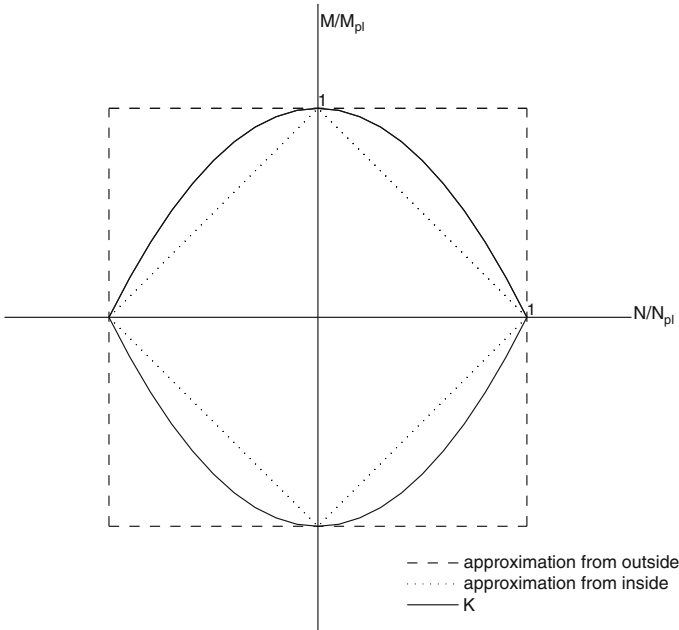


Fig. 8.1 Domain $K_{0,sym}$ with possible approximations

$$\left(\frac{t_i}{N_{ipl}}\right)^2 + \left|\frac{m_i^-}{M_{ipl}}\right| \leq 1, \quad (8.19a)$$

$$\left(\frac{t_i}{N_{ipl}}\right)^2 + \left|\frac{m_i^+}{M_{ipl}}\right| \leq 1. \quad (8.19b)$$

Remark 8.6. Because of the connection between the normal force t_i and the bending moments, (8.19a,b) is also called “ M – N -interaction”.

If the M – N -interaction is not taken into account, $K_{0,\text{sym}}$ is approximated from outside, see Fig. 8.1, by

$$K_{0,\text{sym}}^u := \{(x, y)^T : |x|, |y| \leq 1\}. \quad (8.20)$$

Hence, (8.17a–d) are replaced, cf. (8.19a,b), by the simpler conditions

$$|t_i| \leq N_{ipl} \quad (8.21a)$$

$$|m_i^-| \leq M_{ipl} \quad (8.21b)$$

$$|m_i^+| \leq M_{ipl}. \quad (8.21c)$$

Since the symmetry condition (8.16a) does not hold in general, some modifications of the basic conditions (8.19a,b) are needed. In the non symmetric case $K_{0,xm}$ must be replaced by the intersection

$$K_0 = K_0^U \cap K_0^L \quad (8.22)$$

of two convex sets K_0^U and K_0^L . For a rectangular cross-sectional area we have

$$K_0^U = \{(x, y)^T : x \leq \sqrt{1 - |y|}, |y| \leq 1\} \quad (8.23a)$$

for tension and

$$K_0^L = \{(x, y)^T : -x \leq \sqrt{1 - |y|}, |y| \leq 1\} \quad (8.23b)$$

compression, where again $x = \frac{N}{N_{ipl}}$ and $y = \frac{M}{M_{ipl}}$ (see Fig. 8.2).

In case of tension, see (8.17b,d), from (8.23a) we obtain then the feasibility condition

$$\frac{t_i}{N_{ipl}^U} \leq \sqrt{1 - \left|\frac{m_i^-}{M_{ipl}}\right|} \quad (8.24a)$$

$$\frac{t_i}{N_{ipl}^U} \leq \sqrt{1 - \left|\frac{m_i^+}{M_{ipl}}\right|} \quad (8.24b)$$

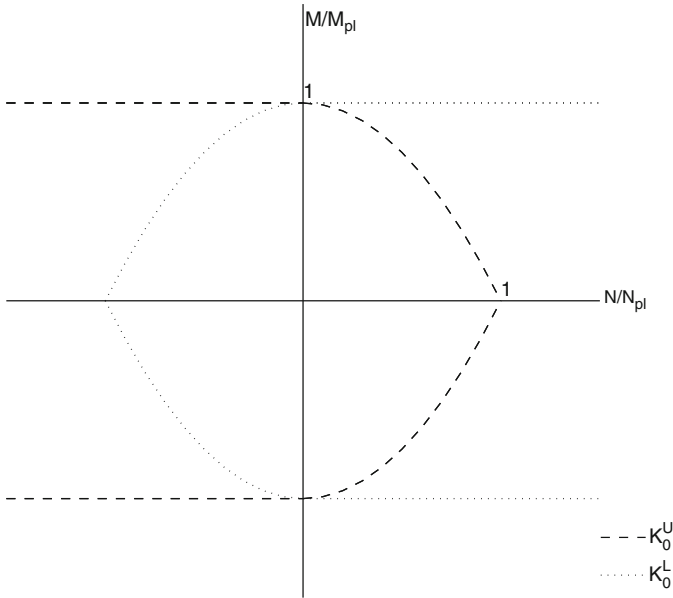


Fig. 8.2 Feasible domain as intersection of K_0^U and K_0^L

$$\left| \frac{m_i^-}{M_{i\,pl}} \right| \leq 1 \tag{8.24c}$$

$$\left| \frac{m_i^+}{M_{i\,pl}} \right| \leq 1. \tag{8.24d}$$

For compression, with (8.17a,c) and (8.23b) we get the feasibility condition

$$-\frac{t_i}{N_{i\,pl}^L} \leq \sqrt{1 - \left| \frac{m_i^-}{M_{i\,pl}} \right|} \tag{8.24e}$$

$$-\frac{t_i}{N_{i\,pl}^L} \leq \sqrt{1 - \left| \frac{m_i^+}{M_{i\,pl}} \right|} \tag{8.24f}$$

$$\left| \frac{m_i^-}{M_{i\,pl}} \right| \leq 1 \tag{8.24g}$$

$$\left| \frac{m_i^+}{M_{i\,pl}} \right| \leq 1. \tag{8.24h}$$

From (8.24a), (8.24e) we get

$$-N_{ipl}^L \sqrt{1 - \left| \frac{m_i^-}{M_{ipl}} \right|} \leq t_i \leq N_{ipl}^U \sqrt{1 - \left| \frac{m_i^-}{M_{ipl}} \right|}. \quad (8.25a)$$

and (8.24b), (8.24f) yield

$$-N_{ipl}^L \sqrt{1 - \left| \frac{m_i^+}{M_{ipl}} \right|} \leq t_i \leq N_{ipl}^U \sqrt{1 - \left| \frac{m_i^+}{M_{ipl}} \right|}. \quad (8.25b)$$

Furthermore, (8.24c), (8.24g) and (8.24d), (8.24h) yield

$$|m_i^-| \leq M_{ipl} \quad (8.25c)$$

$$|m_i^+| \leq M_{ipl}. \quad (8.25d)$$

For computational purposes, piecewise linearizations are applied [30] to the nonlinear conditions (8.25a,b), see also [3, 15]. A basic approximation of K_0^L and K_0^U is given by

$$K_0^{Uu} := \{(x, y)^T : x \leq 1, |y| \leq 1\} =]\infty, 1] \times [-1, 1] \quad (8.26a)$$

and

$$K_0^{Lu} := \{(x, y)^T : x \geq -1, |y| \leq 1\} = [-1, \infty] \times [-1, 1] \quad (8.26b)$$

with $x = \frac{N}{N_{ipl}}$ and $y = \frac{M}{M_{ipl}}$ (see Fig. 8.3).

Since in this approximation the M - N -interaction is not taken into account, condition (8.17a–d) is reduced to

$$-N_{ipl}^L \leq t_i \leq N_{ipl}^U \quad (8.27a)$$

$$|m_i^-| \leq M_{ipl} \quad (8.27b)$$

$$|m_i^+| \leq M_{ipl}. \quad (8.27c)$$

8.2.1 Yield Condition in Case of $M - N$ -Interaction

8.2.1.1 Symmetric Yield Stresses

Consider first the case

$$\sigma_{yi}^U = -\sigma_{yi}^L =: \sigma_{yi}, \quad i = 1, \dots, B. \quad (8.28a)$$

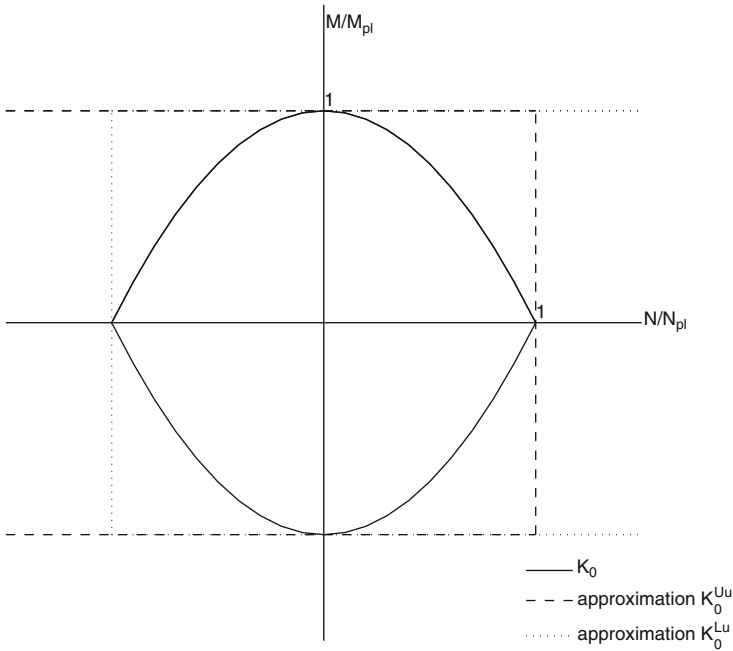


Fig. 8.3 Approximation of K_0 by K_0^{Lu} and K_0^{Lu}

Then,

$$N_{ipl}^L := |\sigma_{yi}^L| A_i = \sigma_{yi}^U A_i =: N_{ipl}^U, \tag{8.28b}$$

hence,

$$N_{ipl} := N_{ipl}^L = N_{ipl}^U = \sigma_{yi} A_i. \tag{8.28c}$$

Moreover,

$$M_{ipl} = \sigma_{yi}^U W_{ipl} = \sigma_{yi} W_{ipl} = \sigma_{yi} A_i \bar{y}_{ic}, \quad i = 1, \dots, B, \tag{8.28d}$$

where \bar{y}_{ic} denotes the arithmetic mean of the centroids of the two half areas of the cross-sectional area A_i of bar i .

Depending on the geometric form of the cross-sectional areal (rectangle, circle, etc.), for the element load vectors

$$F_i = \begin{pmatrix} t_i \\ m_i^+ \\ m_i^- \end{pmatrix}, \quad i = 1, \dots, B, \tag{8.29}$$

of the bars we have the yield condition:

$$\left| \frac{t_i}{N_{ipl}} \right|^\alpha + \left| \frac{m_i^-}{M_{ipl}} \right| \leq 1 \quad (\text{negative end}) \quad (8.30a)$$

$$\left| \frac{t_i}{N_{ipl}} \right|^\alpha + \left| \frac{m_i^+}{M_{ipl}} \right| \leq 1 \quad (\text{positive end}). \quad (8.30b)$$

Here, $\alpha > 1$ is a constant depending on the type of the cross-sectional area of the i th bar. Defining the convex set

$$K_0^\alpha := \left\{ \begin{pmatrix} x \\ y \end{pmatrix} \in \mathbb{R}^2 : |x|^\alpha + |y| \leq 1 \right\}, \quad (8.31)$$

for (8.30a,b) we have also the representation

$$\begin{pmatrix} \frac{t_i}{N_{ipl}} \\ \frac{m_i^-}{M_{ipl}} \end{pmatrix} \in K_0^\alpha \quad (\text{negative end}) \quad (8.32a)$$

$$\begin{pmatrix} \frac{t_i}{N_{ipl}} \\ \frac{m_i^+}{M_{ipl}} \end{pmatrix} \in K_0^\alpha \quad (\text{positive end}). \quad (8.32b)$$

Piecewise Linearization of K_0^α

Due to the symmetry of K_0^α with respect to the transformation

$$x \rightarrow -x, y \rightarrow -y,$$

K_0^α is piecewise linearized as follows.

Starting from a boundary point of K_0^α , hence,

$$\begin{pmatrix} u_1 \\ u_2 \end{pmatrix} \in \partial K_0^\alpha \quad \text{with} \quad u_1 \geq 0, u_2 \geq 0, \quad (8.33a)$$

we consider the gradient of the boundary curve

$$f(x, y) := |x|^\alpha + |y| - 1 = 0$$

of K_0^α in the four points

$$\begin{pmatrix} u_1 \\ u_2 \end{pmatrix}, \begin{pmatrix} -u_1 \\ u_2 \end{pmatrix}, \begin{pmatrix} -u_1 \\ -u_2 \end{pmatrix}, \begin{pmatrix} u_1 \\ -u_2 \end{pmatrix}. \quad (8.33b)$$

We have

$$\nabla f \begin{pmatrix} u_1 \\ u_2 \end{pmatrix} = \begin{pmatrix} \alpha u_1^{\alpha-1} \\ 1 \end{pmatrix} \quad (8.34a)$$

$$\nabla f \begin{pmatrix} -u_1 \\ u_2 \end{pmatrix} = \begin{pmatrix} -\alpha(-(-u_1))^{\alpha-1} \\ 1 \end{pmatrix} = \begin{pmatrix} -\alpha u_1^{\alpha-1} \\ 1 \end{pmatrix} \quad (8.34b)$$

$$\nabla f \begin{pmatrix} -u_1 \\ -u_2 \end{pmatrix} = \begin{pmatrix} -\alpha(-(-u_1))^{\alpha-1} \\ -1 \end{pmatrix} = \begin{pmatrix} -\alpha u_1^{\alpha-1} \\ -1 \end{pmatrix} \quad (8.34c)$$

$$\nabla f \begin{pmatrix} u_1 \\ -u_2 \end{pmatrix} = \begin{pmatrix} \alpha u_1^{\alpha-1} \\ -1 \end{pmatrix}, \quad (8.34d)$$

where

$$\nabla f \begin{pmatrix} -u_1 \\ -u_2 \end{pmatrix} = -\nabla f \begin{pmatrix} u_1 \\ u_2 \end{pmatrix} \quad (8.35a)$$

$$\nabla f \begin{pmatrix} u_1 \\ -u_2 \end{pmatrix} = -\nabla f \begin{pmatrix} -u_1 \\ u_2 \end{pmatrix}. \quad (8.35b)$$

Furthermore, in the two special points

$$\begin{pmatrix} 0 \\ 1 \end{pmatrix} \quad \text{and} \quad \begin{pmatrix} 0 \\ -1 \end{pmatrix}$$

of ∂K_0^α we have, cf. (8.34a), (8.34d), resp., the gradients

$$\nabla f \begin{pmatrix} 0 \\ 1 \end{pmatrix} = \begin{pmatrix} 0 \\ 1 \end{pmatrix} \quad (8.36a)$$

$$\nabla f \begin{pmatrix} 0 \\ -1 \end{pmatrix} = \begin{pmatrix} 0 \\ -1 \end{pmatrix}. \quad (8.36b)$$

Though $f(x, y) = |x|^\alpha + |y| - 1$ is not differentiable at

$$\begin{pmatrix} 1 \\ 0 \end{pmatrix} \quad \text{and} \quad \begin{pmatrix} -1 \\ 0 \end{pmatrix},$$

we define

$$\nabla f \begin{pmatrix} 1 \\ 0 \end{pmatrix} := \begin{pmatrix} 1 \\ 0 \end{pmatrix} \quad (8.36c)$$

$$\nabla f \begin{pmatrix} -1 \\ 0 \end{pmatrix} := \begin{pmatrix} -1 \\ 0 \end{pmatrix}. \quad (8.36d)$$

Using a boundary point $\begin{pmatrix} u_1 \\ u_2 \end{pmatrix}$ of K_0^α with $u_1, u_2 > 0$, the feasible domain K_0^α can be approximated from outside by the convex polyhedron defined as follows.

From the gradients (8.36a–d) we obtain next to the already known conditions (no $M-N$ -interaction):

$$\begin{aligned} \nabla f \begin{pmatrix} 0 \\ 1 \end{pmatrix}^T \left(\begin{pmatrix} x \\ y \end{pmatrix} - \begin{pmatrix} 0 \\ 1 \end{pmatrix} \right) &= \begin{pmatrix} 0 \\ 1 \end{pmatrix}^T \begin{pmatrix} x \\ y-1 \end{pmatrix} \leq 0 \\ \nabla f \begin{pmatrix} 0 \\ -1 \end{pmatrix}^T \left(\begin{pmatrix} x \\ y \end{pmatrix} - \begin{pmatrix} 0 \\ -1 \end{pmatrix} \right) &= \begin{pmatrix} 0 \\ -1 \end{pmatrix}^T \begin{pmatrix} x \\ y+1 \end{pmatrix} \leq 0 \\ \nabla f \begin{pmatrix} 1 \\ 0 \end{pmatrix}^T \left(\begin{pmatrix} x \\ y \end{pmatrix} - \begin{pmatrix} 1 \\ 0 \end{pmatrix} \right) &= \begin{pmatrix} 1 \\ 0 \end{pmatrix}^T \begin{pmatrix} x-1 \\ y \end{pmatrix} \leq 0 \\ \nabla f \begin{pmatrix} -1 \\ 0 \end{pmatrix}^T \left(\begin{pmatrix} x \\ y \end{pmatrix} - \begin{pmatrix} -1 \\ 0 \end{pmatrix} \right) &= \begin{pmatrix} -1 \\ 0 \end{pmatrix}^T \begin{pmatrix} x+1 \\ y \end{pmatrix} \leq 0. \end{aligned}$$

This yields

$$\begin{aligned} y-1 &\leq 0 \\ -1(y+1) &\leq 0 \\ x-1 &\leq 0 \\ -1(x+1) &\leq 0 \end{aligned}$$

or

$$|x| \leq 1 \quad (8.37a)$$

$$|y| \leq 1. \quad (8.37b)$$

Moreover, with the gradients (8.34a–d), cf. (8.35a,b), we get the additional conditions

$$\begin{aligned} \nabla f \begin{pmatrix} u_1 \\ u_2 \end{pmatrix}^T \left(\begin{pmatrix} x \\ y \end{pmatrix} - \begin{pmatrix} u_1 \\ u_2 \end{pmatrix} \right) \\ = \begin{pmatrix} \alpha u_1^{\alpha-1} \\ 1 \end{pmatrix}^T \begin{pmatrix} x-u_1 \\ y-u_2 \end{pmatrix} \leq 0 \quad (1\text{st quadrant}) \end{aligned} \quad (8.38a)$$

$$\begin{aligned} \nabla f \begin{pmatrix} -u_1 \\ -u_2 \end{pmatrix}^T \left(\begin{pmatrix} x \\ y \end{pmatrix} - \begin{pmatrix} -u_1 \\ -u_2 \end{pmatrix} \right) \\ = - \begin{pmatrix} \alpha u_1^{\alpha-1} \\ 1 \end{pmatrix}^T \begin{pmatrix} x+u_1 \\ y+u_2 \end{pmatrix} \leq 0 \quad (3\text{rd quadrant}) \end{aligned} \quad (8.38b)$$

$$\begin{aligned} \nabla f \begin{pmatrix} -u_1 \\ u_2 \end{pmatrix}^T \left(\begin{pmatrix} x \\ y \end{pmatrix} - \begin{pmatrix} -u_1 \\ u_2 \end{pmatrix} \right) \\ = \begin{pmatrix} -\alpha u_1^{\alpha-1} \\ 1 \end{pmatrix}^T \begin{pmatrix} x + u_1 \\ y - u_2 \end{pmatrix} \leq 0 \quad (2\text{nd quadrant}) \end{aligned} \quad (8.38c)$$

$$\begin{aligned} \nabla f \begin{pmatrix} u_1 \\ -u_2 \end{pmatrix}^T \left(\begin{pmatrix} x \\ y \end{pmatrix} - \begin{pmatrix} u_1 \\ -u_2 \end{pmatrix} \right) \\ = - \begin{pmatrix} -\alpha u_1^{\alpha-1} \\ 1 \end{pmatrix}^T \begin{pmatrix} x - u_1 \\ y + u_2 \end{pmatrix} \leq 0 \quad (4\text{th quadrant}). \end{aligned} \quad (8.38d)$$

This means

$$\alpha u_1^{\alpha-1} x - \alpha u_1^\alpha + y - u_2 = \alpha u_1^{\alpha-1} x + y - (\alpha u_1^\alpha + u_2) \leq 0 \quad (8.39a)$$

$$- (\alpha u_1^{\alpha-1} x + \alpha u_1^\alpha + y + u_2) = - (\alpha u_1^{\alpha-1} x + y + (\alpha u_1^\alpha + u_2) -) \leq 0 \quad (8.39b)$$

$$-\alpha u_1^{\alpha-1} x - \alpha u_1^\alpha + y - u_2 = -\alpha - u_1^{\alpha-1} x + y - (\alpha u_1^\alpha + u_2) \leq 0 \quad (8.39c)$$

$$-\alpha u_1^{\alpha-1} x - \alpha u_1^\alpha - y - u_2 = \alpha - u_1^{\alpha-1} x - y - (\alpha u_1^\alpha + u_2) \leq 0. \quad (8.39d)$$

With

$$\alpha u_1^\alpha + u_2 = \nabla f \begin{pmatrix} u_1 \\ u_2 \end{pmatrix}^T \begin{pmatrix} u_1 \\ u_2 \end{pmatrix} =: \beta(u_1, u_2) \quad (8.40)$$

we get the equivalent constraints

$$\begin{aligned} \alpha u_1^{\alpha-1} x + y - \beta(u_1, u_2) &\leq 0 \\ \alpha u_1^{\alpha-1} x + y + \beta(u_1, u_2) &\geq 0 \\ -\alpha u_1^{\alpha-1} x + y - \beta(u_1, u_2) &\leq 0 \\ \alpha u_1^{\alpha-1} x - y - \beta(u_1, u_2) &\leq 0. \end{aligned}$$

This yields the double inequalities

$$|\alpha u_1^{\alpha-1} x + y| \leq \beta(u_1, u_2) \quad (8.41a)$$

$$|\alpha u_1^{\alpha-1} x - y| \leq \beta(u_1, u_2). \quad (8.41b)$$

Thus, a point $u = \begin{pmatrix} u_1 \\ u_2 \end{pmatrix} \in \partial K_0^\alpha$, $u_1 > 0, u_2 > 0$, generates therefore the inequalities

$$-1 \leq x \leq 1 \quad (8.42a)$$

$$-1 \leq y \leq 1 \quad (8.42b)$$

$$-\beta(u_1, u_2) \leq \alpha u_1^{\alpha-1} x + y \leq \beta(u_1, u_2) \quad (8.42c)$$

$$-\beta(u_1, u_2) \leq \alpha u_1^{\alpha-1} x - y \leq \beta(u_1, u_2). \quad (8.42d)$$

Obviously, each further point $\hat{u} \in \partial K_0^\alpha$ with $\hat{u}_1 > 0, \hat{u}_2 > 0$ yields additional inequalities of the type (8.42c,d).

Condition (8.42a–d) can be represented in the following vectorial form:

$$-\begin{pmatrix} 1 \\ 1 \end{pmatrix} \leq I \begin{pmatrix} x \\ y \end{pmatrix} \leq \begin{pmatrix} 1 \\ 1 \end{pmatrix} \quad (8.43a)$$

$$-\beta(u_1, u_2) \begin{pmatrix} 1 \\ 1 \end{pmatrix} \leq H(u_1, u_2) \begin{pmatrix} x \\ y \end{pmatrix} \leq \beta(u_1, u_2) \begin{pmatrix} 1 \\ 1 \end{pmatrix}, \quad (8.43b)$$

with the matrices

$$I = \begin{pmatrix} 1 & 0 \\ 0 & 1 \end{pmatrix}, \quad H(u_1, u_2) = \begin{pmatrix} \alpha u_1^{\alpha-1} & 1 \\ \alpha u_1^{\alpha-1} & -1 \end{pmatrix}. \quad (8.44)$$

Choosing a further boundary point \hat{u} of K_0^α with $\hat{u}_1 > 0, \hat{u}_2 > 0$, we get additional conditions of the type (8.43b).

Using (8.42a–d), for the original yield condition (8.32a,b) we get then the approximative feasibility condition:

1. Negative end of the bar

$$-N_{ipl} \leq t_i \leq N_{ipl} \quad (8.45a)$$

$$-M_{ipl} \leq m_i^- \leq M_{ipl} \quad (8.45b)$$

$$-\beta(u_1, u_2) \leq \alpha u_1^{\alpha-1} \frac{t_i}{N_{ipl}} + \frac{m_i^-}{M_{ipl}} \leq \beta(u_1, u_2) \quad (8.45c)$$

$$-\beta(u_1, u_2) \leq \alpha u_1^{\alpha-1} \frac{t_i}{N_{ipl}} - \frac{m_i^-}{M_{ipl}} \leq \beta(u_1, u_2). \quad (8.45d)$$

2. Positive end of the bar

$$-N_{ipl} \leq t_i \leq N_{ipl} \quad (8.45e)$$

$$-M_{ipl} \leq m_i^+ \leq M_{ipl} \quad (8.45f)$$

$$-\beta(u_1, u_2) \leq \alpha u_1^{\alpha-1} \frac{t_i}{N_{ipl}} + \frac{m_i^+}{M_{ipl}} \leq \beta(u_1, u_2) \quad (8.45g)$$

$$-\beta(u_1, u_2) \leq \alpha u_1^{\alpha-1} \frac{t_i}{N_{ipl}} - \frac{m_i^+}{M_{ipl}} \leq \beta(u_1, u_2). \quad (8.45h)$$

Defining

$$\Gamma^{(i)} := \begin{pmatrix} \frac{1}{N_{ipl}} & 0 & 0 \\ 0 & \frac{1}{M_{ipl}} & 0 \\ 0 & 0 & \frac{1}{M_{ipl}} \end{pmatrix}, \quad F_i = \begin{pmatrix} t_i \\ m_i^+ \\ m_i^- \end{pmatrix}, \quad (8.46)$$

conditions (8.45a–h) can be represented also by

$$-\begin{pmatrix} 1 \\ 1 \\ 1 \end{pmatrix} \leq \Gamma^{(i)} F_i \leq \begin{pmatrix} 1 \\ 1 \\ 1 \end{pmatrix} \quad (8.47a)$$

$$-\beta(u_1, u_2) \begin{pmatrix} 1 \\ 1 \\ 1 \\ 1 \end{pmatrix} \leq \begin{pmatrix} \alpha u_1^{\alpha-1} & 1 & 0 \\ \alpha u_1^{\alpha-1} & -1 & 0 \\ \alpha u_1^{\alpha-1} & 0 & 1 \\ \alpha u_1^{\alpha-1} & 0 & -1 \end{pmatrix} \Gamma_i F_i \leq \beta(u_1, u_2) \begin{pmatrix} 1 \\ 1 \\ 1 \\ 1 \end{pmatrix}. \quad (8.47b)$$

Multiplying (8.45a,c,d,g,h) with N_{ipl} , due to

$$\frac{N_{ipl}}{M_{ipl}} = \frac{\sigma_{yi} A_i}{\sigma_{yi} W_{ipl}} = \frac{\sigma_{yi} A_i}{\sigma_{yi} A_i \bar{y}_{ic}} = \frac{1}{\bar{y}_{ic}}, \quad (8.48)$$

for (8.45a,c,d,g,h) we also have

$$-\beta(u_1, u_2) N_{ipl} \leq \alpha u_1^{\alpha-1} t_i + \frac{m_i^-}{\bar{y}_{ic}} \leq \beta(u_1, u_2) N_{ipl} \quad (8.49a)$$

$$-\beta(u_1, u_2) N_{ipl} \leq \alpha u_1^{\alpha-1} t_i - \frac{m_i^-}{\bar{y}_{ic}} \leq \beta(u_1, u_2) N_{ipl} \quad (8.49b)$$

$$-\beta(u_1, u_2) N_{ipl} \leq \alpha u_1^{\alpha-1} t_i + \frac{m_i^+}{\bar{y}_{ic}} \leq \beta(u_1, u_2) N_{ipl} \quad (8.49c)$$

$$-\beta(u_1, u_2) N_{ipl} \leq \alpha u_1^{\alpha-1} t_i - \frac{m_i^+}{\bar{y}_{ic}} \leq \beta(u_1, u_2) N_{ipl}. \quad (8.49d)$$

8.2.2 Approximation of the Yield Condition by Using Reference Capacities

According to (8.31), (8.32a,b) for each bar $i = 1, \dots, B$ we have the condition

$$\left(\frac{t}{N_{ipl}}, \frac{m}{M_{ipl}} \right) \in K_0^\alpha = \left\{ \begin{pmatrix} x \\ y \end{pmatrix} \in \mathbb{R}^2 : |x|^\alpha + |y| \leq 1 \right\}$$

with $(t, m) = (t_i, m_i^\pm)$, $(N_{ipl}, M_{ipl}) = (N_{ipl}, \bar{M}_{ipl})$.

Selecting fixed reference capacities

$$N_{i0} > 0, M_{i0} > 0, i = 1, \dots, B,$$

related to the plastic capacities N_{ipl}, M_{ipl} , we get

$$\left| \frac{t_i}{N_{i\text{pl}}} \right|^\alpha + \left| \frac{m_i^\pm}{M_{i\text{pl}}} \right| = \left| \frac{t_i}{N_{i0}} \cdot \frac{1}{\frac{N_{i\text{pl}}}{N_{i0}}} \right|^\alpha + \left| \frac{m_i^\pm}{M_{i0}} \cdot \frac{1}{\frac{M_{i\text{pl}}}{M_{i0}}} \right|.$$

Putting

$$\rho_i = \rho_i(a(\omega), x) := \min \left\{ \frac{N_{i\text{pl}}}{N_{i0}}, \frac{M_{i\text{pl}}}{M_{i0}} \right\}, \quad (8.50)$$

we have

$$\frac{\rho_i}{\frac{N_{i\text{pl}}}{N_{i0}}} \leq 1, \quad \frac{\rho_i}{\frac{M_{i\text{pl}}}{M_{i0}}} \leq 1$$

and therefore

$$\left| \frac{t_i}{N_{i\text{pl}}} \right|^\alpha + \left| \frac{m_i^\pm}{M_{i\text{pl}}} \right| = \left| \frac{t_i}{\rho_i N_{i0}} \right|^\alpha \cdot \left| \frac{\rho_i}{\frac{N_{i\text{pl}}}{N_{i0}}} \right|^\alpha + \left| \frac{m_i^\pm}{\rho_i M_{i0}} \right| \cdot \left| \frac{\rho_i}{\frac{M_{i\text{pl}}}{M_{i0}}} \right| \leq \left| \frac{t_i}{\rho_i N_{i0}} \right|^\alpha + \left| \frac{m_i^\pm}{\rho_i M_{i0}} \right|. \quad (8.51)$$

Thus, the yield condition (8.30a,b) or (8.32a,b) is guaranteed by

$$\left| \frac{t_i}{\rho_i N_{i0}} \right|^\alpha + \left| \frac{m_i^\pm}{\rho_i M_{i0}} \right| \leq 1$$

or

$$\left(\frac{\frac{t_i}{\rho_i N_{i0}}}{\frac{m_i^\pm}{\rho_i M_{i0}}} \right) \in K_0^\alpha. \quad (8.52)$$

Applying the piecewise linearization described in Sect. 8.2.1 to condition (8.52), we obtain, cf. (8.45a–h), the approximation stated below. Of course, conditions (8.45a,b,e,f) are not influenced by this procedure. Thus, we find

$$-N_{i\text{pl}} \leq t_i \leq N_{i\text{pl}} \quad (8.53a)$$

$$-M_{i\text{pl}} \leq m_i^- \leq M_{i\text{pl}} \quad (8.53b)$$

$$-M_{i\text{pl}} \leq m_i^+ \leq M_{i\text{pl}} \quad (8.53c)$$

$$-\beta(u_1, u_2) \leq \alpha u_1^{\alpha-1} \frac{t_i}{\rho_i N_{i0}} + \frac{m_i^-}{\rho_i M_{i0}} \leq \beta(u_1, u_2) \quad (8.53d)$$

$$-\beta(u_1, u_2) \leq \alpha u_1^{\alpha-1} \frac{t_i}{\rho_i N_{i0}} - \frac{m_i^-}{\rho_i M_{i0}} \leq \beta(u_1, u_2) \quad (8.53e)$$

$$-\beta(u_1, u_2) \leq \alpha u_1^{\alpha-1} \frac{t_i}{\rho_i N_{i0}} + \frac{m_i^+}{\rho_i M_{i0}} \leq \beta(u_1, u_2) \quad (8.53f)$$

$$-\beta(u_1, u_2) \leq \alpha u_1^{\alpha-1} \frac{t_i}{\rho_i N_{i0}} - \frac{m_i^+}{\rho_i M_{i0}} \leq \beta(u_1, u_2). \quad (8.53g)$$

Remark 8.7. Multiplying with ρ_i we get quotients $\frac{t_i}{N_{i0}}, \frac{m_i^\pm}{M_{i0}}$ with fixed denominators.

Hence, multiplying (8.53d–g) with ρ_i , we get the equivalent system

$$-N_{ipl} \leq t_i \leq N_{ipl} \quad (8.54a)$$

$$-M_{ipl} \leq m_i^- \leq M_{ipl} \quad (8.54b)$$

$$-M_{ipl} \leq m_i^+ \leq M_{ipl} \quad (8.54c)$$

$$-\beta(u_1, u_2)\rho_i \leq \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} + \frac{m_i^-}{M_{i0}} \leq \beta(u_1, u_2)\rho_i \quad (8.54d)$$

$$-\beta(u_1, u_2)\rho_i \leq \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} - \frac{m_i^-}{M_{i0}} \leq \beta(u_1, u_2)\rho_i \quad (8.54e)$$

$$-\beta(u_1, u_2)\rho_i \leq \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} + \frac{m_i^+}{M_{i0}} \leq \beta(u_1, u_2)\rho_i \quad (8.54f)$$

$$-\beta(u_1, u_2)\rho_i \leq \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} - \frac{m_i^+}{M_{i0}} \leq \beta(u_1, u_2)\rho_i. \quad (8.54g)$$

Obviously, (8.54a–g) can be represented also in the following form:

$$|t_i| \leq N_{ipl} \quad (8.55a)$$

$$|m_i^-| \leq M_{ipl} \quad (8.55b)$$

$$|m_i^+| \leq M_{ipl} \quad (8.55c)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} + \frac{m_i^-}{M_{i0}} \right| \leq \beta(u_1, u_2)\rho_i \quad (8.55d)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} - \frac{m_i^-}{M_{i0}} \right| \leq \beta(u_1, u_2)\rho_i \quad (8.55e)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} + \frac{m_i^+}{M_{i0}} \right| \leq \beta(u_1, u_2)\rho_i \quad (8.55f)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} - \frac{m_i^+}{M_{i0}} \right| \leq \beta(u_1, u_2)\rho_i. \quad (8.55g)$$

By means of definition (8.50) of ρ_i , system (8.55a–g) reads

$$|t_i| \leq N_{ipl} \quad (8.56a)$$

$$|m_i^-| \leq M_{ipl} \quad (8.56b)$$

$$|m_i^+| \leq M_{ipl} \quad (8.56c)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} + \frac{m_i^-}{M_{i0}} \right| \leq \beta(u_1, u_2) \frac{N_{ipl}}{N_{i0}} \quad (8.56d)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} - \frac{m_i^-}{M_{i0}} \right| \leq \beta(u_1, u_2) \frac{M_{ipl}}{M_{i0}} \quad (8.56e)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} - \frac{m_i^-}{M_{i0}} \right| \leq \beta(u_1, u_2) \frac{N_{i pl}}{N_{i0}} \quad (8.56f)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} - \frac{m_i^-}{M_{i0}} \right| \leq \beta(u_1, u_2) \frac{M_{i pl}}{M_{i0}} \quad (8.56g)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} + \frac{m_i^+}{M_{i0}} \right| \leq \beta(u_1, u_2) \frac{N_{i pl}}{N_{i0}} \quad (8.56h)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} + \frac{m_i^+}{M_{i0}} \right| \leq \beta(u_1, u_2) \frac{M_{i pl}}{M_{i0}} \quad (8.56i)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} - \frac{m_i^+}{M_{i0}} \right| \leq \beta(u_1, u_2) \frac{N_{i pl}}{N_{i0}} \quad (8.56j)$$

$$\left| \alpha u_1^{\alpha-1} \frac{t_i}{N_{i0}} - \frac{m_i^+}{M_{i0}} \right| \leq \beta(u_1, u_2) \frac{M_{i pl}}{M_{i0}} \quad (8.56k)$$

Corresponding to Remark 8.7, the variables

$$t_i, m_i^+, m_i^-, A_i \text{ or } x$$

enters linearly. Increasing the accuracy of approximation by taking a further point $\hat{u} = (\hat{u}_1, \hat{u}_2)$ with the related points $(\hat{u}_1, -\hat{u}_2)$, $(-\hat{u}_1, \hat{u}_2)$, $(-\hat{u}_1, -\hat{u}_2)$, we obtain further inequalities of the type (8.55d–g), (8.56d–g) respectively.

8.3 Stochastic Optimization

Due to (8.1c), (8.10), (8.11c), (8.12c), the $3B$ – vector

$$F = (F_1^T, \dots, F_B^T)^T \quad (8.57a)$$

of all interior loads fulfills the equilibrium condition

$$CF = R \quad (8.57b)$$

with the external load vector R and the equilibrium matrix C .

In the following we collect all random model parameters [31, 35, 39], such as external load factors, material strength parameters, cost factors, etc., into the random v -vector

$$a = a(\omega), \quad \omega \in (\Omega, \mathcal{A}, \mathcal{P}), \quad (8.58a)$$

on a certain probability space (Ω, \mathcal{A}, P) .

Thus, since in some cases the vector R of external loads depend also on the design r -vector x , we get

$$R = R(a(\omega), x). \quad (8.58b)$$

Of course, the plastic capacities depend also on the vectors x and $a(\omega)$, hence,

$$N_{ipl}^{\Gamma} = N_{ipl}^{\Gamma}(a(\omega), x), \quad \Gamma = L, U \quad (8.58c)$$

$$M_{ipl} = M_{ipl}(a(\omega), x). \quad (8.58d)$$

We assume that the probability distribution and/or the needed moments of the random parameter vector $a = a(\omega)$ are known [2, 28, 31, 42].

The remaining deterministic constraints for the design vector x are represented by

$$x \in \mathcal{D} \quad (8.59)$$

with a certain convex subset D of \mathbb{R}^r .

8.3.1 Violation of the Yield Condition

Consider in the following an interior load distribution F fulfilling the equilibrium condition (8.58b).

According to the analysis given in Sect. 8.2, after piecewise linearization, the yield condition for the i th bar can be represented by an inequality of the type

$$H\Gamma^{(i)}(a(\omega), x) \leq h^{(i)}(a(\omega), x), \quad i = 1, \dots, B, \quad (8.60)$$

with matrices $H, \Gamma^{(i)} = \Gamma^{(i)}(a(\omega), x)$ and a vector $h^{(i)} = h^{(i)}(a(\omega), x)$ as described in Sect. 8.2.

In order to take into account violations of condition (8.60), we consider the equalities

$$H\Gamma^{(i)}F_i + z_i = h^{(i)}, \quad i = 1, \dots, B. \quad (8.61)$$

If

$$z_i \geq 0, \quad \text{for all } i = 1, \dots, B, \quad (8.62a)$$

then (8.60) holds, and the yield condition is then fulfilled too, or holds with a prescribed accuracy.

However, in case

$$z_i \not\geq 0 \quad \text{for some bars } i \in \{1, \dots, B\}, \quad (8.62b)$$

the survival condition is violated at some points of the structure. Hence, structural failures may occur. The resulting costs Q of failure, damage and reconstructure of the frame is a function of the vectors $z_i, i = 1, \dots, B$, defined by (8.61). Thus, we have

$$Q = Q(z) = Q(z_1, \dots, z_B), \quad (8.63a)$$

where

$$z := (z_1^T, z_2^T, \dots, z_B^T)^T, \quad (8.63b)$$

$$z_i := h^{(i)} - H\Gamma^{(i)}F_i, \quad i = 1, \dots, B. \quad (8.63c)$$

8.3.2 Cost Function

Due to the survival condition (8.62a), we may consider cost functions Q such that

$$Q(z) = 0, \quad \text{if } z \geq 0, \quad (8.64)$$

hence, no (recourse) costs arise if the yield condition (8.60) holds.

In many cases the recourse or failure costs of the structure are defined by the sum

$$Q(z) = \sum_{i=1}^B Q_i(z_i), \quad (8.65)$$

of the element failure costs $Q_i = Q_i(z_i), i = 1, \dots, B$.

Using the representation

$$z_i = y_i^+ - y_i^-, \quad y_i^+, y_i^- \geq 0, \quad (8.66a)$$

the member cost functions $Q_i = Q_i(z_i)$ are often defined [20, 31] by considering the linear function

$$q_i^{-T} y_i^- + q_i^{+T} y_i^+ \quad (8.66b)$$

with certain vectors q_i^+, q_i^- of cost coefficients for the evaluation of the condition $z_i \geq 0, z_i \not\geq 0$, respectively.

The cost function $Q_i = Q_i(z_i)$ is then defined by the minimization problem

$$\min q_i^{-T} y_i^- + q_i^{+T} y_i^+ \quad (8.67a)$$

$$\text{s.t. } y_i^+ - y_i^- = z_i \quad (8.67b)$$

$$y_i^-, y_i^+ \geq 0. \quad (8.67c)$$

If $z_i := (z_{i1}, \dots, z_{i\mu})^T, y_i^\pm := (y_{i1}^\pm, \dots, y_{i\mu}^\pm)^T, i = 1, \dots, B$ then (8.67a–c) can also be represented by

$$\min \sum_{l=1}^{\mu} (q_{il}^- y_{il}^- + q_{il}^+ y_{il}^+) \quad (8.68a)$$

$$\text{s.t. } y_{il}^+ - y_{il}^- = z_{il}, \quad l = 1, \dots, \mu \quad (8.68b)$$

$$y_{il}^-, y_{il}^+ \geq 0, \quad l = 1, \dots, \mu. \quad (8.68c)$$

Since the pairs of variables (y_{il}^-, y_{il}^+) , $l = 1, \dots, \mu$, are not connected with each other by the constraints, and the objective function is separable with respect to these pairs of variables, (8.68a–c) can be decomposed into μ separated minimization problems

$$\min q_{il}^- y_{il}^- + q_{il}^+ y_{il}^+ \quad (8.69a)$$

$$\text{s.t. } y_{il}^+ - y_{il}^- = z_{il} \quad (8.69b)$$

$$y_{il}^-, y_{il}^+ \geq 0, \quad (8.69c)$$

for the pairs of variables (y_{il}^-, y_{il}^+) , $l = 1, \dots, \mu$.

Under the condition

$$q_{il}^- + q_{il}^+ \geq 0, l = 1, \dots, \mu, \quad (8.70)$$

the following result holds:

Lemma 8.1. *Suppose that (8.70) holds. Then the minimum value function $Q_{il} = Q_{il}(z_{il})$ of (8.69a–c) is a piecewise linear, convex function given by*

$$Q_{il}(z_{il}) := \max\{q_{il}^+ z_{il}, -q_{il}^- z_{il}\}. \quad (8.71)$$

Hence, the member cost functions $Q_i = Q_i(z_i)$ reads

$$Q_i(z_i) = \sum_{l=1}^{\mu} Q_{il}(z_{il}) = \sum_{l=1}^{\mu} \max\{q_{il}^+ z_{il}, -q_{il}^- z_{il}\}, \quad (8.72a)$$

and the total cost function $Q = Q(z)$ is given by

$$Q(z) = \sum_{i=1}^B Q_i(z_i) = \sum_{i=1}^B \sum_{l=1}^{\mu} \max\{q_{il}^+ z_{il}, -q_{il}^- z_{il}\}. \quad (8.72b)$$

8.3.3 Choice of the Cost Factors

Under elastic conditions the change $\Delta\sigma$ of the total stress σ and the change ΔL of the element length L are related by

$$\Delta L = \frac{L}{E} \Delta\sigma, \quad (8.73a)$$

where E denotes the modulus of elasticity.

Assuming that the neutral axis is equal to the axis of symmetry of the element, for the total stress $\Delta\sigma$ in the upper (“+”) lower (“-”) fibre of the boundary we have

$$\Delta\sigma = \frac{\Delta t}{A} \pm \frac{\Delta m}{W}, \quad (8.73b)$$

where W denotes the axial modulus of the cross-sectional area of the element (beam).

Representing the change ΔV of volume of an element by

$$\Delta V = A \cdot \Delta L, \quad (8.74a)$$

then

$$\begin{aligned} \Delta V &= A \cdot \Delta L = A \cdot \frac{L}{E} \Delta\sigma = A \cdot \frac{L}{E} \left(\frac{\Delta t}{A} \pm \frac{\Delta m}{W} \right) \\ &= \frac{L}{E} \Delta t \pm \frac{L}{E} \frac{A}{W} \Delta m = \frac{L}{E} \Delta t \pm \frac{L}{E} \frac{1}{\bar{y}_c} \Delta m = \frac{L}{E} \Delta t \pm \frac{L}{E} \cdot \frac{1}{\bar{y}_c} \Delta m, \end{aligned} \quad (8.74b)$$

where \bar{y}_c is the cross-sectional parameter as defined in (8.28d).

Consequently, due to (8.73a,b), for the evaluation of violations Δt of the axial force constraint we may use a cost factor of the type

$$\Gamma_K := \frac{L}{E}, \quad (8.75a)$$

and an appropriate cost factor for the evaluation of violations of moment constraints reads

$$\Gamma_M = \frac{L}{E} \cdot \frac{1}{\bar{y}_c}. \quad (8.75b)$$

8.3.4 Total Costs

Denoting by

$$G_0 = G_0(a(\omega), x) \quad (8.76a)$$

the primary costs, such as weighted negative load factors, material costs, costs of construction, etc., the total costs including failure or recourse costs are given by

$$G = G_0(a(\omega), x) + Q(z(a(\omega), x, F(\omega))). \quad (8.76b)$$

Hence, the total costs $G = G(a(\omega), x, F(\omega))$ depend on the vector $x = (x_1, \dots, x_r)^T$ of design variables, the random vector $a(\omega) = (a_1(\omega), \dots, a_v(\omega))^T$ of model parameters and the random vector $F = F(\omega)$ of all internal loadings.

Minimizing the expected total costs, we get the following stochastic optimization problem (SOP) of recourse type [20]

$$\min E \left(G_0(a(\omega), x) + Q(z(a(\omega), x, F(\omega))) \right) \tag{8.77a}$$

$$\text{s.t. } H\Gamma^{(i)}(a(\omega), x)F_i(\omega) + z_i(\omega) = h^{(i)}(a(\omega), x) \quad \text{a.s.,} \\ i = 1, \dots, B \tag{8.77b}$$

$$CF(\omega) = R(a(\omega), x) \quad \text{a.s.} \tag{8.77c}$$

$$x \in D. \tag{8.77d}$$

Using representation (8.65), (8.67a–c) of the recourse or failure cost function $Q = Q(z)$, problem (8.77a–d) takes also the following equivalent from

$$\min E \left(G_0(a(\omega), x) + \sum_{i=1}^B (q_i^-(\omega)^T y_i^-(\omega) + q_i^+(\omega)^T y_i^+(\omega)) \right) \tag{8.78a}$$

$$\text{s.t. } H\Gamma^{(i)}(a(\omega), x)F_i(\omega) + y_i^+(\omega) - y_i^-(\omega) = h^{(i)}(a(\omega), x) \quad \text{a.s.,} \\ i = 1, \dots, B \tag{8.78b}$$

$$CF(\omega) = R(a(\omega), x) \quad \text{a.s.} \tag{8.78c}$$

$$x \in D, y_i^+(\omega), y_i^-(\omega) \geq 0 \quad \text{a.s., } i = 1, \dots, B. \tag{8.78d}$$

Remark 8.8. Stochastic optimization problems of the type (8.78a–d) are called “two-stage stochastic programs” or “stochastic problems with recourse”.

In many cases the primary cost function $G_0 = G_0(a(\omega), x)$ represents the volume or weight of the structural, hence,

$$\begin{aligned} G_0(a(\omega), x) &:= \sum_{i=1}^B \Gamma_i(\omega)V_i(x) \\ &= \sum_{i=1}^B \Gamma_i(\omega)L_i A_i(x), \end{aligned} \tag{8.79}$$

with certain (random) weight factors $\Gamma_i = \Gamma_i(\omega)$.

In case

$$A_i(x) := u_i x_i \tag{8.80}$$

with fixed sizing parameters $u_i, i = 1, \dots, B$, we get

$$\begin{aligned} G_0(a(\omega), x) &= \sum_{i=1}^B \Gamma_i(\omega)L_i u_i(x) = \sum_{i=1}^B \Gamma_i(\omega)L_i h_i x_i \\ &= c(a(\omega))^T x, \end{aligned} \tag{8.81a}$$

where

$$c(a(\omega)) := (\Gamma_1(\omega)L_1u_1, \dots, \Gamma_B(\omega)L_Bu_B)^T. \quad (8.81b)$$

Thus, in case (8.80), $G_0 = G_0(a(\omega), x)$ is a linear function of x .

8.3.5 Discretization Methods

The expectation in the objective function of the stochastic optimization problem (8.78a–d) must be computed numerically. One of the main methods is based on the discretization of the probability distribution $P_{a(\cdot)}$ of the random parameter ν -vector $a = a(\omega)$, hence,

$$P_{a(\cdot)} \approx \mu := \sum_{k=1}^s \alpha_k \epsilon_{a^{(k)}} \quad (8.82a)$$

with

$$\alpha_k \geq 0, \quad k = 1, \dots, s, \quad \sum_{k=1}^s \alpha_k = 1. \quad (8.82b)$$

Corresponding to the realizations $a^{(k)}$, $k = 1, \dots, s$, of the discrete approximate (8.82a,b), we have the realizations $y_i^{-(k)}$, $y_i^{+(k)}$ and $F^{(k)}$, $F_i^{(k)}$, $k = 1, \dots, s$, of the random vectors $y_i^-(\omega)$, $y_i^+(\omega)$, $i = 1, \dots, B$, and $F(\omega)$. then

$$\begin{aligned} & E \left(G_0(a(\omega), x) + \sum_{i=1}^B (q_i^{-T} y_i^-(\omega) + q_i^{+T} y_i^+(\omega)) \right) \\ & \approx \sum_{k=1}^s \alpha_k \left(G_0(a^{(k)}, x) + \sum_{i=1}^B (q_i^{-T} y_i^{-(k)} + q_i^{+T} y_i^{+(k)}) \right). \end{aligned} \quad (8.83a)$$

Furthermore, the equilibrium equation (8.77c) is approximated by

$$CF^{(k)} = R(a^{(k)}, x), \quad k = 1, \dots, s, \quad (8.83b)$$

where $F^{(k)} := (F_1^{(k)T}, \dots, F_B^{(k)T})^T$, and we have, cf. (8.78d), the nonnegativity constraints

$$y_i^{+(k)}, y_i^{-(k)} \geq 0, \quad k = 1, \dots, s, \quad i = 1, \dots, B. \quad (8.83c)$$

Thus, (SOP) (8.78a–d) is reduced to the parameter optimization problem

$$\min \bar{G}_0(a^{(k)}, x) + \sum_{i=1}^B \alpha_k (q_i^{-T} y_i^{-(k)} + q_i^{+T} y_i^{+(k)}) \quad (8.84a)$$

$$\text{s.t. } H\Gamma^{(i)}(a^{(k)}, x)F_i^{(k)} + y_i^{+(k)} - y_i^{-(k)} = h^{(i)}(a^{(k)}, x),$$

$$i = 1, \dots, B, k = 1, \dots, s \quad (8.84b)$$

$$CF^{(k)} = R(a^{(k)}, x), k = 1, \dots, s \quad (8.84c)$$

$$x \in D, y_i^{+(k)}, y_i^{-(k)} \geq 0, k = 1, \dots, s, i = 1, \dots, B. \quad (8.84d)$$

A further important class of methods for computing expectations and probabilities, hence, multiple integrals, occurring in stochastic optimization problems, reliability analysis and reliability-based optimal design (RBO), cf. [9, 11, 30, 31], are simulation methods, such as Monte Carlo Simulation (MCS) procedures. Simulation techniques are used especially in cases with only few information about the analytical properties of the underlying technical device, e.g., in case of analytically almost unavailable limit state functions. In principle, MSC is a very simple technique which is widely applicable on the one hand, but may have a very low efficiency of estimation on the other hand. Hence, several improvements were considered in the last time, such as Advanced Monte Carlo Simulation techniques: Variance reduction methods reducing the sampling error, based, e.g., on importance sampling methods, direction sampling, subset simulation, etc., see, e.g., [4]. Further improvements can be obtained by combining these simulation/estimation techniques with Response Surface Methods (RSM) for estimating unknown functions using regression techniques and advanced nonlinear programming procedures (NLP), cf. [12].

8.3.6 Complete Recourse

According to Sect. 8.3.1, the evaluation of the violation of the yield condition (8.60) is based on (8.61), hence

$$H\Gamma^{(i)}F_i + z_i = h^{(i)}, i = 1, \dots, B.$$

In generalization of the so-called “simple recourse” case (8.67a–c), in the “complete recourse” case the deviation

$$z_i = h^{(i)} - H\Gamma^{(i)}F_i$$

is evaluated by means of the minimum value $Q_i = Q_i(z_i)$ of the linear program, cf. (8.68a–c)

$$\min q^{(i)T} y^{(i)} \quad (8.85a)$$

$$\text{s.t. } M^{(i)} y^{(i)} = z_i \quad (8.85b)$$

$$y^{(i)} \geq 0. \quad (8.85c)$$

Here, $q^{(i)}$ is a given cost vector and $M^{(i)}$ denotes the so-called recourse matrix [20].

We assume that the linear equation (8.85b) has a solution $y^{(i)} \geq 0$ for each vector z_i . This property is called “complete recourse”.

In the present case the stochastic optimization problem (8.77a–c) reads

$$\min E \left(G_0(a(\omega), x) + \sum_{i=1}^B q^{(i)T} y^{(i)}(\omega) \right) \quad (8.86a)$$

$$\text{s.t. } H\Gamma^{(i)}(a(\omega), x) F_i(\omega) + M^{(i)} y^{(i)}(\omega) = h^{(i)}(a(\omega), x) \text{ a.s.,} \\ i = 1, \dots, B \quad (8.86b)$$

$$CF(\omega) = R(a(\omega), x) \text{ a.s.} \quad (8.86c)$$

$$x \in D, y^{(i)}(\omega) \geq 0 \text{ a.s., } i = 1, \dots, B. \quad (8.86d)$$

As described in Sect. 8.3.5, problem (8.86a–d) can be solved numerically by means of discretization methods and application of linear/nonlinear programming techniques.

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Part IV
Analysis and Optimization of Economic
and Engineering Systems Under
Uncertainty

Chapter 9

Uncertainty in the Future Nitrogen Load to the Baltic Sea Due to Uncertain Meteorological Conditions

Jerzy Bartnicki

Abstract The Norwegian Meteorological Institute has a long-term project with HELCOM Commission for regular calculation of annual atmospheric deposition of nitrogen to the Baltic Sea. In 2005, the institute received an additional project from HELCOM with the aim of estimating atmospheric nitrogen deposition to six sub-basins and catchments of the Baltic Sea for the year 2010, using nitrogen emission projections according to agreed emission ceilings under the EU National Emission Ceilings (NEC) Directive and the Gothenburg Protocol. Since, the meteorology for 2010 is unknown, model calculations were performed for four selected years with different meteorology: 1996, 1997, 1998 and 2000, which are available in the database. Final deposition values for the year 2010 were calculated as an average over the four selected years. In this way we were able to estimate the uncertainty restricted to meteorological variability. The ranges between minimum and maximum of calculated depositions to sub-basins and catchments are large indicating significant variation of the deposition depending on meteorological conditions.

9.1 Introduction

The Helsinki Commission, or HELCOM is the governing body of the “Convention on the Protection of the Marine Environment of the Baltic Sea Area” – known as the Helsinki Convention [7]. The important role of HELCOM is to protect the marine environment of the Baltic Sea from all sources of pollution through intergovernmental co-operation between Denmark, Estonia, the European Community, Finland, Germany, Latvia, Lithuania, Poland, Russia and Sweden, which are the Contracting Parties to HELCOM. HELCOM’s vision for the future is a healthy Baltic Sea environment with diverse biological components functioning in balance, resulting in a good ecological status and supporting a wide range of sustainable economic and social activities (<http://www.helcom.fi/>).

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Eutrophication is one of the major problems for the Baltic Sea. Since the 1800s, the Baltic Sea has changed from an oligotrophic clear-water sea into a eutrophic marine environment. Nitrogen and phosphorus are among the main growth-limiting nutrients and as such do not pose any direct hazards to marine organisms. Eutrophication, however, is a condition in an aquatic ecosystem where high nutrient concentrations stimulate growth of algae which leads to imbalanced functioning of the system. It is mainly caused by a significant nutrient load to the Baltic Sea (<http://www.helcom.fi/>).

The nutrient inputs entering the Baltic Sea are either airborne or waterborne. The main pathways of nutrient input to the Baltic Sea are:

- Direct atmospheric deposition on the Baltic Sea water surface
- Riverine inputs of nutrients to the sea. Rivers transport nutrients that have been discharged or lost to inland surface waters within the Baltic Sea catchment area
- Point sources discharging directly to the sea.

Atmospheric deposition of nitrogen accounts for approximately 30% of the total nitrogen load to the Baltic Sea. Therefore, nitrogen has been regularly monitored by analysing the results of measurements and model calculations in co-operation with the Co-operative Programme for Monitoring and Evaluation of the Long-range Transmission of Air pollutants in Europe (EMEP). The main objective of the EMEP is to regularly provide Governments and subsidiary bodies under the Convention on Long-range Transboundary Air Pollution [8] with qualified scientific information to support the development and further evaluation of the international protocols on emission reductions negotiated under the Convention (<http://www.emep.int/>). A co-operation between EMEP and HELCOM was established already in 1996. Since then, three EMEP Centres have published joint reports estimating annual supply of nitrogen heavy metals and persistent organic pollutants to the Baltic Sea. Updated emissions, as well as results of measurements and their analysis are also included in the annual EMEP reports for HELCOM. Such an annual report has been recently prepared for HELCOM in 2007 [1].

In 2004 HELCOM established a project using models to assess the implications of different policy scenarios on nutrient inputs and the resulting eutrophication status in order to indicate the most cost-effective measures for the different sub-regions of the Baltic Sea. Most of the models considered under HELCOM have been ecological models related to the assessment of effects to the sea. The basis of the effect models were the scenarios of activities on land. Therefore, the aim was to link management scenario models with ecological models in order to assess measures for reducing nutrient inputs. The aim of the project was to assess for HELCOM Contracting Parties the impact of different agricultural policy scenarios on nutrient inputs in the Baltic Sea catchment area and on the eutrophication status of the Baltic Sea. The final aim was to enable the identification of cost-effective measures in the different parts of the Baltic Sea catchment area which requires achieving a good ecological status throughout the Baltic Sea area [2].

In 2005 HELCOM considered airborne nitrogen pollution and decided to include nitrogen air depositions into the ongoing HELCOM Project. To evaluate the

implications of different policy scenarios on nutrient inputs, HELCOM agreed that EMEP should be charged with the task of assessing the changes in atmospheric nitrogen deposition under the condition that nitrogen emission targets according to the Gothenburg protocol to the LRTAP Convention and the EU NEC Directive as projected for 2010 are fulfilled. Based on the above decision, EMEP has received a project from HELCOM.

As requested by HELCOM, all nitrogen depositions as well as source allocation budgets have been calculated for the six sub-basins and catchments of the Baltic Sea for the year 2010 using the EMEP Unified model [10] (<http://www.emep.int/>). Names and acronyms of the sub-basins and catchments used in the model calculations are given below:

1. Gulf of Bothnia (GUB)
2. Gulf of Finland (GUF)
3. Gulf of Riga (GUR)
4. Baltic Proper (BAP)
5. Belt Sea (BES)
6. The Kattegat (KAT)

Depositions and source allocation budgets have been also calculated for the entire basin and the entire catchment of the Baltic Sea. Geographical borders of sub-basins and catchments used in the computations are shown in Fig. 9.1.

Emissions and meteorological fields are the main input into the EMEP Unified model which calculates transport and deposition of air pollutants over Europe. For the historical calculations, meteorological fields are available from the Numerical Weather Prediction model HIRLAM [3] which is run operationally at the Norwegian Meteorological Institute in Oslo. Since the meteorological fields are not known for 2010, we have used the meteorological fields from the past years, which are



Fig. 9.1 Locations of six sub-basins and catchments of the Baltic Sea for which the nitrogen depositions for 2010 have been calculated

available in the EMEP database. Variability between different meteorological years in the past create significant uncertainty in the model results for 2010. Analysis of this specific uncertainty is the main subject of the current paper.

9.2 Nitrogen Emissions

Emission input to the EMEP Unified model requires information about annual nitrogen oxides and ammonia emissions from all sources located in the EMEP domain.

Projections for 2010 assume nitrogen oxides and ammonia national emissions as specified in the NEC Directive [9] for selected 15 EU countries. National nitrogen oxides and ammonia emissions as specified in the Gothenburg Protocol [4] are assumed for all countries listed therein except for those already mentioned in the NEC Directive. Emissions for the Russian Federation and Estonia are taken from the HELCOM publication [6]. Finally, 2010 projections of nitrogen emissions due to international ship traffic on the Europeans seas are taken from the EMEP database following Entec projections [5]. All remaining emission sources of nitrogen are taken from the EMEP database <http://www.emep.int/>.

9.2.1 National Emission Ceilings According to EU NEC Directive

2010 national emission ceilings for nitrogen oxides (units: kt of NO₂ per year) and ammonia (units: kt of NH₃ per year) according to the EU NEC Directive [9] are shown in Table 9.1.

The national emission ceilings presented in Table 9.1 were designed with the aim of broadly meeting the interim environmental objectives set out in Article 5 of the Directive. It is supposed that the Community area with depositions of nutrient nitrogen in excess of the critical loads will be reduced by about 30% compared to 1990.

Table 9.1 2010 national emission ceilings for nitrogen oxides (units: kt of NO₂ per year) and for ammonia (units: kt of NH₃ per year) according to the EU NEC Directive [9]

Country	NO _x	NH ₃	Country	NO _x	NH ₃
Austria	103	66	Belgium	176	74
Denmark	127	69	Finland	170	31
France	810	780	Germany	1,051	550
Greece	44	73	Ireland	65	116
Italy	90	419	Luxembourg	11	7
Netherlands	260	128	Portugal	250	90
Spain	847	353	Sweden	148	57
United Kingdom	1,167	297	EC15	6,519	3,110

Table 9.2 2010 national emission ceilings of nitrogen oxides (units: kt of NO₂ per year) and ammonia (units: kt of NH₃ per year) according to the 1999 Gothenburg Protocol [4]

Country	NO _x	NH ₃	Country	NO _x	NH ₃
Armenia	46	25	Austria	107	66
Belarus	255	158	Belgium	181	74
Bulgaria	266	108	Croatia	87	30
Czech Republic	286	101	Denmark	127	69
Finland	170	31	France	860	780
Germany	1,081	550	Greece	344	73
Hungary	198	90	Ireland	65	116
Italy	1,000	419	Latvia	84	44
Liechtenstein	0.37	0.15	Lithuania	110	84
Luxembourg	11	7	Netherlands	266	128
Norway	156	23	Poland	879	468
Portugal	260	108	Republic of Moldova	90	42
Romania	437	210	Russian Federation (PEMA)	265	49
Slovakia	130	39	Slovenia	45	20
Spain	847	353	Sweden	148	57
Switzerland	79	63	Ukraine	1,222	592
United Kingdom	1,181	297	European Community	6,671	3,129

Among the EU-15 countries, United Kingdom (1,167 kt NO₂) and Germany (1,051 kt NO₂) are the main NO₂ emitters in 2010. France (780 kt NH₃) and Germany (550 kt NH₃) are the main NH₃ sources.

During the EU accession process national emission ceilings were reduced for Latvia to 61 kt NO₂. This most recent value for Latvia [6] and the nitrogen emissions shown in Table 9.2 were used for the EMEP model run. The same emission sources are also considered in the Gothenburg Protocol, however NEC emissions of nitrogen oxides and ammonia projected for 2010 are lower or the same as those projected by the Gothenburg Protocol presented in the next section.

9.2.2 National Emission Ceilings According to Gothenburg Protocol

The national emission ceilings for nitrogen oxides and ammonia specified in the Protocol to Abate Acidification, Eutrophication and Ground-level Ozone done in Gothenburg, Sweden on 30 November 1999 [4] are shown in Table 9.2.

For the Russian Federation the Protocol specifies only the emission ceilings for the so-called Pollutant Emissions Management Area (PEMA). In the EMEP domain, 2010 nitrogen oxide and ammonia emissions for the entire Russian Federation are 2,653 kt NO₂ and 1,179 kt NH₃, respectively.

Compared to emission levels in 1990, main reductions of nitrogen oxides emissions should occur in the Czech Republic (61%), Germany (60%), Sweden

(56%) and United Kingdom (56%). Main reductions of ammonia emissions in 2010 compared to 1990 are expected in Denmark (43%), Netherlands (43%), Slovakia (37%) and Czech Republic (35%).

9.2.3 Nitrogen Emission Projections Used in the Model Runs

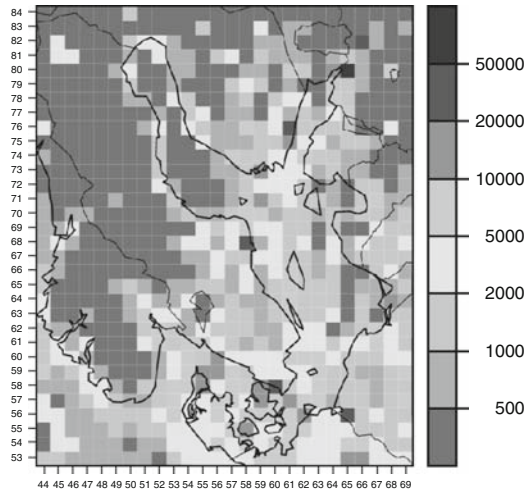
Projected national annual total emissions of nitrogen oxides and ammonia used in the model runs are shown in Table 9.3 for 2010. All anthropogenic sources in the EMEP area are taken into account in Table 9.3, including emissions from the HELCOM parties and from the international ship traffic on the Baltic Sea.

The numbers in Table 9.3 are based on total emission targets for each country, including areas outside the Baltic Sea Catchment, which explains high values for, e.g. Russia and Germany. Not all emissions in the Contracting Parties end up in the Baltic Sea. For 2010, the total nitrogen oxide emissions target in the EMEP area is approximately 19.7 million tones of NO₂, whereas the total ammonia emissions target in the EMEP area is about 7.3 million tones of NH₃.

Table 9.3 2010 national total emissions of nitrogen oxides (units: kt of NO₂ per year) and ammonia (units: kt of NH₃ per year) used in the model runs

Source	NO _x	NH ₃	Source	NO _x	NH ₃
Albania	27.9	25.9	Armenia	46.0	25.0
Austria	102.9	66.2	Azerbaijan	43.0	25.0
Baltic Sea	457.5	0.0	Belarus	254.9	157.8
Belgium	176.1	74.4	Black Sea	154.8	0.0
Bosnia and Herz.	54.0	17.3	Bulgaria	265.5	107.6
Croatia	97.3	30.1	Cyprus	21.1	6.3
Czech Republic	286.1	101.4	Denmark	126.7	69.3
Estonia	59.0	28.4	Finland	170.3	30.8
France	810.3	780.3	Georgia	30.0	97.0
Germany	1,051.2	549.8	Greece	344.5	72.5
Hungary	198.4	89.5	Iceland	30.0	3.0
Ireland	65.0	115.8	Italy	990.1	418.88
Kazakhstan	50.2	19.0	Latvia	60.8	43.8
Lithuania	110.4	84.4	Luxembourg	11.0	7.1
Macedonia	5.9	1.4	Mediterranean Sea	2,382	80.0
Moldova, Republic of	90.6	41.6	Netherlands	260.3	128.3
North Africa	96.0	235.0	North Sea	862.4	0.0
Norway	156.0	23.0	Norway	156.0	23.0
Poland	878.3	467.9	Remaining Asiatic ar.	9.0	178.0
Remaining NE Atlantic	740.5	0.0	Romania	436.5	209.7
Russian Federation	2,653.0	1,178.8	Serbia and Montenegro	167.8	69.3
Slovakia	130.2	19.9	Spain	148.4	57.3
Switzerland	79.3	62.6	Turkey	851.8	240.6
Ukraine	1,222.2	591.9	United Kingdom	1,167.4	296.6
EMEP	19,686.3	7,345.2			

Fig. 9.2 Spatial distribution of nitrogen oxide emissions around the Baltic Sea for 2010, used in the EMEP model computations. Units: tonnes of NO₂ per year and per 50 km × 50 km grid cell



The total nitrogen oxides emissions and ammonia emissions targets in the HELCOM Parties are estimated to be 5.3 million tones of NO₂ and 2.5 million tones of NH₃, respectively.

Table 9.3 exhibits that the Russian Federation and ship traffic on the Mediterranean Sea are the largest emitters of nitrogen oxides in 2010, with 2,653 and 2,383 kt NO₂, respectively. These two sources are approximately twice as large than the next on the list: Ukraine (1,222 kt NO₂), United Kingdom (1,167 kt NO₂), Germany (1,051 kt NO₂) and Italy (990 kt NO₂).

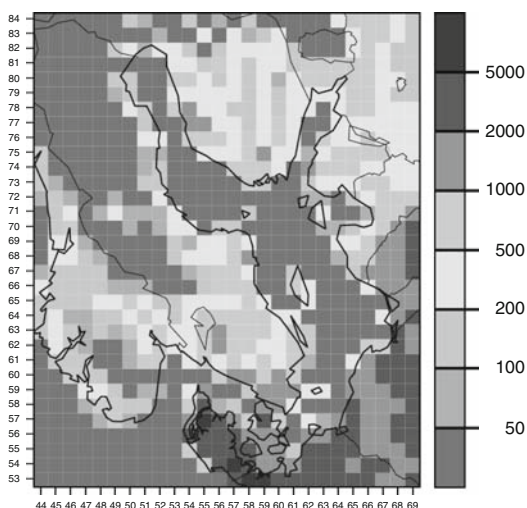
The Russian Federation (1,179 kt NH₃), France (780 kt NH₃), Ukraine (592 kt NH₃) and Germany (550 kt NH₃) belong to the four highest emitters of ammonia in 2010. The emission targets for the rest of the countries are much lower with Poland (550 kt NH₃) and Italy (468 kt NH₃) taking the fifth and sixth places, respectively.

To run the EMEP model, the emissions should be distributed in the model grid system. Maps with the 2010 nitrogen oxides and ammonia emissions in 2010 spatially distributed over the Baltic Sea region are shown in Figs.9.2 and 9.3 respectively.

9.3 Computed Nitrogen Depositions for 2010

Nitrogen emission inventories described in the previous chapter have been used in order to estimate nitrogen depositions in 2010. Since, the meteorology for 2010 is unknown, model calculations had to be performed with existing meteorological data from the past years. Meteorological data for the years 1996, 1997, 1998 and 2000 and emission projections for the year 2010 have been used for the EMEP Unified model runs. Final deposition values for the year 2010 were calculated as the average over the four selected years.

Fig. 9.3 Spatial distribution of ammonia emissions around the Baltic Sea for 2010, used in the EMEP model computations. Units: tonnes of NH_3 per year and per $50 \text{ km} \times 50 \text{ km}$ grid cell



9.3.1 Unified EMEP Model

The Unified EMEP model is an Eulerian model that has been developed at the Meteorological Synthesizing Centre West of EMEP (EMEP/MSC-W) for simulating atmospheric transport and deposition of acidifying and eutrophying compounds as well as photo-oxidants in Europe. The model has been documented in the EMEP Status Report [10]. Here we only give a short description of the basic features of the model. Model details as well as recent changes and updates can be found on the EMEP web site <http://www.emep.int/>.

The model domain covers Europe and the Atlantic Ocean. The model grid (of the size 170×133) has a horizontal resolution of 50 km at 60°N , which is consistent with the resolution of emission data reported to EMEP. In the vertical, the model has 20 sigma layers reaching up to 100 hPa . Approximately 10 of these layers are placed below 2 km to obtain a high resolution for the boundary layer which is of special importance for the long range transport of air pollution.

The EMEP unified model uses 3-hourly resolved meteorological data from the PARLAM-PS model, a dedicated version of the HIRLAM (High Resolution Limited Area Model) Numerical Weather Prediction model operational at the Norwegian Meteorological Institute [3].

The emission input consists of gridded national annual emissions of sulphur dioxide, nitrogen oxides, ammonia, non-methane volatile organic compounds (VOC) and carbon monoxide. They are available in each of the $50 \text{ km} \times 50 \text{ km}$ model grid cells. These emissions are distributed temporally according to monthly and daily factors derived from data provided by the University of Stuttgart (IER).

Concentrations of 71 species are computed in the Unified EMEP model (56 are advected, 15 are short-lived and not advected). The sulphur and nitrogen chemistry is coupled to the photo-chemistry, which allows a more sophisticated description of, e.g. the oxidation of sulphur dioxide to sulphate.

Dry deposition is calculated using the resistance analogy and is a function of the pollutant type, meteorological conditions and surface properties. Parameterization of wet deposition processes includes both in-cloud and sub-cloud scavenging of gases and particles using scavenging coefficients.

9.3.2 Calculated Depositions to Sub-basins and Catchments of the Baltic Sea

Calculated nitrogen depositions to sub-basins and catchments of the Baltic Sea are given in Tables 9.4 and 9.5, respectively. Calculated depositions are shown for each sub-basin and catchment, and for four meteorological years used in the computations. The deposition in 2010 was calculated as an average over four meteorological years.

Locations of minimum and maximum deposition are slightly different for different sub-basins and catchments and for different deposition types. However, for most sub-basins and catchments, the lowest deposition values can be observed for the meteorological year 1997 and the highest for the meteorological year 2000.

For all meteorological years and all sub-basins and catchments, wet nitrogen deposition is significantly higher, sometimes two times higher than dry deposition of nitrogen.

Depositions of oxidized and reduced nitrogen to all sub-basins are approximately on the same level, but deposition of oxidized nitrogen is slightly higher than deposition of reduced nitrogen in the north and in the middle (GUB, GUF, GUR and BAP sub-basins). In the south (BES and KAT sub-basins) deposition of reduced nitrogen is higher. Deposition of oxidized nitrogen is also higher than deposition of reduced nitrogen to the entire basin of the Baltic Sea. In general, there is more oxidized nitrogen than reduced nitrogen deposition into sub-basins of the Baltic Sea.

In the case of catchments, oxidized nitrogen deposition is higher than reduced nitrogen deposition for GUB, GUF and KAT catchments. For GUR, BAP, BES catchments and the for entire catchment of the Baltic Sea, the relation between oxidized and reduced nitrogen deposition is opposite. In general, there is more reduced nitrogen than oxidized nitrogen deposition into catchments of the Baltic Sea.

9.4 Uncertainty Due to Meteorological Variability

Inspection of Tables 9.4 and 9.5 gives the first impression of uncertainty restricted to meteorological variability. To assess the variability of computed depositions due to varying meteorological conditions the standard deviation (σ_{N-1}) was calculated for each of the deposition type and each of sub-basin and catchment of the Baltic Sea.

The ranges between minimum and maximum can also indicate the uncertainty of computed depositions due to variable meteorological conditions. This is an

Table 9.4 2010 nitrogen depositions to sub-basins of the Baltic Sea and to the entire basin of the Baltic Sea calculated with the help of the meteorology from four different years and as average over these years. Units: kt N

Sub-basin	Meteo	Oxidized dry	Oxidized wet	Reduced dry	Reduced wet	Total
GUB	1996	6.1	11.8	2.9	8.9	29.7
GUB	1997	5.2	10.2	2.4	6.8	24.6
GUB	1998	6.5	15.0	2.8	11.0	35.3
GUB	2000	6.9	18.7	3.9	15.7	45.1
GUB	Mean	6.2	13.9	3.0	10.6	33.7
GUF	1996	2.9	5.2	1.9	5.4	15.3
GUF	1997	2.5	4.1	1.6	3.7	11.9
GUF	1998	3.0	5.5	2.1	4.9	15.4
GUF	2000	3.1	6.4	2.3	5.5	17.3
GUF	Mean	2.9	5.3	2.0	4.9	15.0
GUR	1996	2.2	3.3	1.8	3.7	11.0
GUR	1997	1.9	3.3	1.4	3.2	9.8
GUR	1998	2.2	3.5	1.8	3.4	10.9
GUR	2000	2.4	3.9	2.0	3.9	12.2
GUR	Mean	2.2	3.5	1.7	3.6	11.0
BAP	1996	23.1	40.1	23.3	39.7	126.2
BAP	1997	19.6	33.3	20.2	31.4	104.4
BAP	1998	22.2	47.1	23.9	43.8	136.9
BAP	2000	22.9	47.1	26.3	44.1	140.4
BAP	Mean	21.9	41.9	23.4	39.7	127.0
BES	1996	2.6	4.9	5.1	6.3	18.9
BES	1997	2.4	3.9	5.6	5.4	17.3
BES	1998	2.5	5.8	5.5	7.8	21.6
BES	2000	2.5	5.8	6.1	7.3	21.7
BES	Mean	2.5	5.1	5.6	6.7	19.9
KAT	1996	2.5	4.4	3.5	5.0	15.4
KAT	1997	2.3	4.1	3.4	4.6	14.4
KAT	1998	2.3	5.4	3.5	5.7	16.9
KAT	2000	2.6	6.5	4.0	6.7	19.8
KAT	Mean	2.4	5.1	3.6	5.5	16.6
BAS	1996	39.4	69.7	38.5	68.9	216.6
BAS	1997	33.9	58.9	34.6	55.0	182.4
BAS	1998	38.7	82.3	39.5	76.5	237.0
BAS	2000	40.3	88.4	44.5	83.2	256.5
BAS	Mean	38.1	74.8	39.3	70.9	223.1

important indicator of uncertainty from the practical point of view in the decision-making process. The relative ranges and relative standard deviations were expressed in per cent of the mean value of the calculated deposition. The results are presented in Tables 9.6 and 9.7 for sub-basins and catchments, respectively.

Table 9.5 2010 nitrogen depositions to six catchments of the Baltic Sea and to the entire catchment of the Baltic Sea calculated with the help of the meteorology from four different years and as average over these years. Units: kt N

Catchment	Meteo	Oxidized dry	Oxidized wet	Reduced dry	Reduced wet	Total
GUB	1996	36	48	14	39	137
GUB	1997	31	44	12	31	118
GUB	1998	34	58	13	44	150
GUB	2000	42	74	19	59	193
GUB	Mean	36	56	15	43	149
GUF	1996	47	70	22	69	207
GUF	1997	41	66	23	63	194
GUF	1998	51	76	27	72	227
GUF	2000	56	82	31	75	244
GUF	Mean	49	74	26	70	218
GUR	1996	23	34	25	46	128
GUR	1997	20	37	23	49	130
GUR	1998	23	37	26	51	136
GUR	2000	25	36	29	46	136
GUR	Mean	23	36	26	48	132
BAP	1996	128	187	198	273	786
BAP	1997	118	178	202	264	761
BAP	1998	132	209	206	293	840
BAP	2000	138	205	225	279	848
BAP	Mean	129	195	208	277	809
BES	1996	7	11	12	15	46
BES	1997	7	9	13	13	42
BES	1998	7	14	13	19	52
BES	2000	7	14	14	18	53
BES	Mean	7	12	13	16	48
KAT	1996	18	19	14	21	72
KAT	1997	16	18	14	20	68
KAT	1998	16	23	14	24	78
KAT	2000	18	30	16	31	96
KAT	Mean	17	23	15	24	78
BAS	1996	258	370	286	463	1,376
BAS	1997	233	353	288	441	1,314
BAS	1998	264	417	300	503	1,483
BAS	2000	286	442	334	507	1,569
BAS	Mean	260	395	302	478	1,436

The relative ranges are roughly twice as high as the standard deviations and these two measures of uncertainty are well correlated for all kinds of nitrogen depositions. The relative ranges can be interpreted as the worst case scenario when the meteorology in the reference year and in the year 2010 gives the most different results in

Table 9.6 Relative ranges and standard deviations (in brackets) of nitrogen depositions to sub-basins of the Baltic Sea and to the entire basin of the Baltic Sea, calculated for 2010 with meteorology from four different years. Units: per cent of mean nitrogen deposition over four years

Sub-basin	Oxidized dry	Oxidized wet	Reduced dry	Reduced wet	Total
GUB	28 (12)	61 (27)	48 (21)	84 (38)	61 (36)
GUF	22 (10)	42 (17)	35 (15)	38 (17)	36 (15)
GUR	24 (10)	18 (8)	38 (16)	19 (8)	22 (9)
BAP	16 (7)	33 (16)	26 (11)	32 (15)	28 (13)
BES	6 (2)	38 (18)	17 (7)	36 (16)	22 (11)
KAT	10 (5)	47 (21)	15 (7)	38 (17)	32 (14)
BAS	17 (8)	39 (18)	25 (10)	40 (17)	33 (14)

Table 9.7 Relative ranges and standard deviations (in brackets) of nitrogen depositions to the catchments of the Baltic Sea and to the entire catchment of the Baltic Sea, calculated for 2010 with meteorology from four different years. Units: per cent of mean nitrogen deposition over four years

Sub-basin	Oxidized dry	Oxidized wet	Reduced dry	Reduced wet	Total
GUB	30 (13)	53 (24)	46 (20)	64 (27)	50 (21)
GUF	31 (13)	22 (10)	36 (16)	17 (7)	23 (10)
GUR	20 (8)	10 (4)	21 (9)	10 (5)	6 (3)
BAP	16 (7)	16 (8)	13 (6)	10 (4)	11 (5)
BES	3 (1)	39 (18)	16 (6)	35 (16)	22 (11)
KAT	16 (8)	52 (24)	15 (7)	45 (20)	35 (15)
BAS	20 (8)	22 (10)	16 (7)	14 (7)	18 (8)

calculated nitrogen deposition. This is the maximum uncertainty estimated from the four years meteorological database. This can be the case if only one meteorological year is taken as a basis for verification of the emission reductions effects in 2010.

The relative standard deviation shows the deviation from the mean deposition over four years. Therefore, the averaging effect is already included in this measure of uncertainty.

For all sub-basins except GUR the uncertainty of computed wet deposition of both oxidized and reduced nitrogen is much higher than the uncertainty of computed dry deposition. In the extreme case of BES sub-basin, uncertainty of computed wet deposition is several times higher than the uncertainty of computed dry deposition for this sub-basin. The reason is a patchy and irregular pattern of precipitation fields, both in space and time.

The largest uncertainty due to meteorological variability can be noticed for GUB (Gulf of Bothnia) sub-basin and catchment with relatively low nitrogen deposition and significant contribution of nitrogen from the long-range transport to the deposition. On the other hand, for BES and KAT sub-basin uncertainty of computed dry oxidized deposition is relatively small, because these sub-basins are located close to the large emissions sources of nitrogen

The uncertainties of the depositions due to variable meteorology are rather large, both for sub-basins and catchments of the Baltic Sea. However, uncertainties of

nitrogen deposition to the catchments are in general lower than the uncertainties of nitrogen deposition to sub-basins of the Baltic Sea. The reason is a larger area of the catchment compared to the area of corresponding sub-basin and therefore more likely compensation of different types of uncertainties for the catchments.

Taking into account that uncertainties in nitrogen deposition caused by changing meteorology are larger than the expected emission reductions, one cannot expect a proportional impact of emission reduction on the deposition for one specific year 2010. For example, 10% reduction of nitrogen emission in all EMEP sources can give 6% increase of nitrogen deposition in the entire basin of the Baltic Sea in 2010, if meteorological conditions are the same as in the year 2000. However, the uncertainty due to variable meteorology is decreasing when more years are included in the verification period for the effects of emission reductions.

9.5 Conclusions

The largest uncertainty due to meteorological variability in computed nitrogen deposition for 2010 can be noticed for GUB (Gulf of Bothnia) sub-basin and catchment with relatively low deposition fluxes.

The relative ranges and deviations of the depositions due to variable meteorology are rather large compared to expected emission reductions, both for sub-basins and catchments of the Baltic Sea. For the deposition of oxidized and reduced nitrogen to sub-basins and to the entire basin of the Baltic Sea there is more uncertainty in wet than in dry deposition. The uncertainty of nitrogen deposition to the catchments are in general lower than the uncertainty of nitrogen deposition to sub-basins of the Baltic Sea.

The main conclusion from the decision making perspective is that even significant efforts in reducing nitrogen emissions do not need to become immediately visible in modelled and measured nitrogen depositions to the Baltic Sea due to inter-annual variability of meteorological conditions. In order to notice significant deposition changes caused by the emission reduction it is necessary to wait at least several years. It is very important to take this type of uncertainty into account in future policy actions concerning the improvement of the Baltic Sea environment (<http://www.helcom.fi/>).

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Chapter 10

Planning Sustainable Agricultural Development Under Risks

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Abstract In this paper we show that explicit treatment of risks and uncertainties is an essential element in planning sustainable agricultural development. Introduction of risks and uncertainties in production models considerably alter strategies for achieving robust outcomes. We discuss stochastic models that may assist to derive optimal agricultural production allocation and expansion within environmental and health risk indicators. Approaches are illustrated with the example of spatially-explicit livestock production allocation in China to 2030.

10.1 Introduction

Global change, economic-demographic and urbanization growth, changing consumption preferences alter the structure of agricultural production systems. In particular, they promote industrial agriculture geared towards making use of economies of scale to produce the highest output at the lowest cost. Although intensification has shown many positive effects, there are significant disadvantages, risks, and costs involved. Undesirable impacts of intensification include environmental pollution, input-intensive mono-cropping, and the marginalization and decline of smallholder farms, causing abandonment of land and migration of rural population to cities. These are further exacerbated by various risks such as climate change and variability, natural catastrophes, market distortions and instabilities.

Alone environmental impacts and health hazards associated with intensive agricultural production have increased awareness and established the need to identify pathways towards sustainable agriculture.

This paper aims to show that adequate accounting and treatment of risks and uncertainties is a necessary condition for planning sustainable agricultural development. Naturally that considerations of risks may considerably alter production and consumption decisions. This fact is illustrated in Sect. 10.2 with a stylized model of

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two agricultural producers characterized by different levels of efficiency and exposure to risks. The example captures, in a nutshell, the features of a geographically detailed and dynamic model for agricultural production planning under risks and uncertainties, as adopted for the analysis of livestock production development in China to 2030 [12, 14].

Now, a growing share of livestock products in China is coming from industrial and specialized enterprises associated with hazardous pollution of the atmosphere, water and soil resources, which becomes a critical environmental issue [17, 25]. Traditional livestock systems represented a natural farming cycle; livestock was kept on grass areas or in confined places close to farmland. Primary sources of feed were grass, feed from fodder crops and other crops, household wastes and crop residues. In these systems, livestock waste and manure were considered valuable sources of nutrients for crop production or for fuel. The manure was recycled efficiently, causing minimal environmental degradation and pollution. With the introduction of large-scale industrial livestock production, especially of pigs and poultry, this closed cycle is collapsing. Intensive livestock production enterprises are located close to meat markets, near urban areas, and in these locations there is much more livestock concentrated than land can support for proper manure recycling.

Geographical allocation of animals and the levels of intensification at which livestock is kept, differently affect the occurrences and spread of livestock diseases. In a sense, increasing specialization and intensification of livestock production is associated with newly emerging diseases (e.g., possibly SARS, avian flu) that can threaten human health.

Concentration of intensive livestock production is an important cause of environmental pollution and health hazards. When coinciding with intensive crop cultivation, the problem of pollution through excess nutrients from livestock operations is further exacerbated by imbalanced fertilizer application. Over-supply of nutrients may lead to toxic nitrate pollution in the water supply and may cause eutrophication of surface water. The trend is alarming and in some locations, without appropriate measures, it may turn irreversible. The analysis in [14] has shown that the development of China livestock production sector cannot just continue along past intensification trends. The goal of this paper is to discuss model-based approaches to guide decisions regarding the inevitable and significant future expansion of livestock production with respect to economic conditions at locations accounting for sustainability and risk indicators. Indicators of sustainability and risks are defined by various interdependent factors including the spatial distribution of people and incomes, the current levels of livestock production and intensification, and the conditions and current use of land resources. Combinations of these factors are used in proposed models to distinguish different locations by the degree of their risk exposure in order to achieve robust solutions.

In Sect. 10.3 we introduce a stochastic spatially explicit and dynamic simulation model used for planning livestock and crop production expansion coherently with projected demand increases to 2030. It allows for spatio-temporal and risk-adjusted analysis of production developments under alternative socio-economic, demographic, and technological scenarios. This allows to address not

only environmental and social concerns, but also investigate innovative policies offering new viable opportunities to farmers, agricultural workers, consumers, and markets. The approach has also been discussed in [13] with alternative scenario settings.

The meaningful specification of risk indicators and constraints to define alternative allocation scenarios is often constrained by the paucity of data at required resolutions. In this case, specific downscaling (disaggregating) and upscaling (aggregating) procedures [11] provide a tool for estimation of dependencies between the geographical factors, constraints, and economic-environmental policy responses. In Sect. 10.3 we analyze the main features of these procedures for spatial production allocation with respect to risks and suitability constraints in locations. Section 10.4 introduces a new stochastic optimization approach for planning production allocation when some of the risks in the model of Sect. 10.3 are explicitly taken into account by stochastic scenarios. In fact, the Sect. 10.3 and Sect. 10.4 distinguish two types of uncertainties: endogenous uncertainties associated with behavioral principles regarding production expansion and exogenous uncertainties associated with parameters of models. Section 10.5 describes alternative allocation scenarios and presents selected numerical results. Section 10.6 concludes and indicates directions for future work.

10.2 Cooperation and Co-existence for Risk Sharing

Over the last 20 years, China's demand and production of livestock products has increased remarkably due to rapid development of the national economy, urbanization, rising living standards, and population growth [5]. Increasing incomes and changing consumption preferences have boosted production and have shifted the composition of producers towards specialized enterprises with a number of advantages: they are more feed efficient and profitable, flexible in terms of management, may better adjust and comply to legislation, and, in general benefit from economies of scale. In a sense, these trends follow the Ricardo's assertion [24] that trading nations gain from production specialization and intensification. Accordingly, we may expect that production should be undertaken by the most efficient agent, with intensified production on large farms. This is true only under idealized conditions when risks are not accounted for.

In reality, agricultural production facilities may be exposed to various risks, but also may cause different negative impacts. Depending on the location and intensity, values of the facilities are interdependent subject to contingencies, and are determined endogenously. For this reason, of particular interest are production chains with large and small units to stabilize the aggregate production. Contrary to Ricardo, the less efficient and intensive producer may provide the supply of production and enhance market stability, say, if the producer's risks are different and weakly or even negatively correlated with others. Such diversification of producers by scale and location hedges against economic and environmental risks, improves

welfare and ensures continuous supply of agricultural products to markets. Explicit accounting of risks may considerably alter the composition of production units and their intensification levels in a chain.

Let us illustrate this with a stylized model of only two producers, $i = 1, 2$, which in Sect. 10.4 will be extended to a multi-producer case. Let x_i denote the production level of i -th producer and assume that only one good is produced, e.g., meat; c_i is the cost per unit of produce. The product can also be imported from an external source with price b per unit of produce. Assume $c_1 < c_2 < b$, i.e., the cheapest source is the first producer. The production has to satisfy the exogenous inelastic demand d of a given region.

In the *absence of risks*, the model is formulated as the minimization of the total cost function:

$$c_1x_1 + c_2x_2 \quad (10.1)$$

subject to

$$\begin{aligned} x_1 + x_2 &= d, \\ x_1 \geq 0, x_2 &\geq 0, \end{aligned} \quad (10.2)$$

where x_1, x_2 are production capacities. The optimal solution to the problem is $x_1^* = d, x_2^* = 0$, i.e., the production is undertaken by the more efficient producer, which accords with Ricardo's views.

In case of *risk exposure*, the endogenous supply (10.2) is expressed, for example, as a linear function

$$a_1x_1 + a_2x_2 = d, \quad (10.3)$$

where a_1, a_2 are contingencies or "supply" shocks to x_1, x_2 , e.g., due to outbreaks of diseases, weather risks, or other hazardous events. We assume that a_1, a_2 are random variables $0 \leq a_i \leq 1$, which may reduce the supply from $i = 1, 2$. If endogenous supply $a_1x_1 + a_2x_2$ falls short of demand d , the residual amount $d - a_1x_1 - a_2x_2$ must be obtained from external sources at unit import cost b . The planning of production capacities x_1, x_2 can be evaluated from the minimization of total production costs and potential import cost, i.e., the minimization of the function

$$F(x) = c_1x_1 + c_2x_2 + bE \max\{0, d - a_1x_1 - a_2x_2\},$$

where $x_1 \geq 0, x_2 \geq 0$ and the expected import cost when the demand d exceeds the supply $a_1x_1 + a_2x_2$ is $bE \max\{0, d - a_1x_1 - a_2x_2\}$. In this case, the role of a less efficient producer for stabilizing supply is clearly visible.

Assume that only the efficient producer is at risk, that is $a_2 = 1$. Let function $F(x)$ have continuous derivatives, e.g., the cumulative distribution function of a_1 has a continuous density function. It is easy to see that the optimal positive decisions $x_1^* > 0, x_2^* > 0$ exist in the case when partial derivatives meet $F_{x_1}(0, 0) < 0, F_{x_2}(0, 0) < 0$. We have $F_{x_1}(0, 0) = c_1 - bEa_1, F_{x_2}(0, 0) = c_2 - b$ and, perhaps counter intuitively, the less efficient, but without risks producer 2 is active unconditionally (since $c_2 - b < 0$). The cost efficient producer 1 is inactive in the case $c_1 - bEa_1 \geq 0$, leaving production entirely to the higher-cost producer 2. Only in

the case $c_1 - bEa_1 < 0$ both producers are active. Hence, in this example the less cost-efficient producer is able to stabilize the aggregate production in the presence of contingencies affecting the more cost-effective producer 1.

To derive the market share of the producer 2, take the derivative

$$F_{x_2}(x, x_2) = c_2 - bP[d > a_1x + x_2]$$

according to optimality conditions of stochastic minimax problems [8]. This means that the optimal production level $x_2^* > 0$ of producer 2 is a quantile defined by the equation $P[d > a_1x_1^* + x_2^*] = c_2/b$, assuming $x_1^* > 0$ (otherwise $x_2^* = d$). It also depends on x_1^* and all conditions ensuring a positive share x_1^* of producer 1. Although not at risk ($a_2 = 1$), the optimal production level of producer 2 is defined by (10.3) through interdependencies among producers participating in the same market with demand d . Let us now consider the case when both producers are at risks, i.e., $a_1 \neq 1$, $a_2 \neq 1$. The existence of optimal positive production of both producers follows from similar equations

$$F_{x_1}(0, 0) = c_1 - bEa_1 < 0,$$

$$F_{x_2}(0, 0) = c_2 - bEa_2 < 0.$$

The structure of optimal solution is similar to the case when only one producer is at risk. In particular, there may be a situation where $c_2 - bEa_2 \geq 0$, when producer 2 is inactive, but the cost effective producer 1 is active now with the insurance provided by the external source (import or borrowing).

Apart from exogenous risks, the production and the market are subject to endogenous risks dependent on the level of x_1, x_2 . Negative impacts of production increase and intensification cause contamination of water, soil, air in the densely populated areas, which may incur uncertain, possibly highly non-linear costs, increasing with increasing x_1, x_2 . In this case, the cooperation and market sharing may be unconditionally advantageous, as the following case illustrates.

Let us now consider a case when costs are increasing non-linear functions, for the sake of simplicity, *quadratic*, $c_1x_1^2 + c_2x_2^2$, $c_1 < c_2$, and there are no production distortions, i.e., $a_1 = a_2 = 1$. The problem is to minimize

$$c_1x_1^2 + c_2x_2^2$$

subject to the demand-supply constraints

$$x_1 + x_2 = d, x_1 \geq 0, x_2 \geq 0.$$

At least one producer must be active, say producer 1. It is easy to see from the standard optimality condition that the optimal level is $x_1^* = \frac{c_1}{c_1+c_2}d$. Therefore, the optimal level for producer 2 is $x_2^* = \frac{c_2}{c_1+c_2}d$. In other words, both $x_1^* > 0$, $x_2^* > 0$, i.e., unconditional on the cost effectiveness of the producer 1, the increasing

non-linear production costs require co-existence and cooperation of both producers. These examples emphasize that market shares are to a larger extent determined by the production costs, the import price, and the contingencies of producers. In fact, the less efficient but with lower risk, producer will likely have a higher share than a more efficient, but with higher risk exposure, producer. For the sake of simplicity, in the above examples the contingencies are characterized by a probability distribution. In reality, the contingencies, e.g., livestock diseases, environmental pollution, demand fluctuations, economic instabilities, have complex geographical and temporal patterns of occurrences, are subject to spatial interactions. Their mutual probability distribution functions may not be analytically tractable and thus require stochastic simulation models as presented in Sect. 10.3 and downscaling procedures allowing for estimation of required values based on all available auxiliary statistics and model-derived results. Risk exposures are often characterized by certain standards commonly imposed as additional safety constraints on admissible values of some indicators, e.g., constraints on ambient standards in the pollution control.

10.3 Agricultural Planning Under Risks

10.3.1 A Simulation Model

The stochastic and dynamic livestock and crop production model developed by the Land Use Change and Agriculture (LUC) program at the International Institute for Applied Systems Analysis (IIASA) [10, 12] integrates demographic, economic, agricultural and environmental modeling components. The IIASA model is essentially an accounting GIS-based model, which allows to incorporate inherent processes in an endogenized manner.

The model is developed with the aim to assist in planning sustainable agricultural developments combining various national, subnational and regional interacting agricultural activities, production, processing, consumers. Together with reasonable scales of biophysical modeling, this allows for production planning within limited resources and possibilities to improve or recover production potentials, against uncertainties of weather, climate change, market situation or other risks such as the contamination of land or pasture. Simplicity of model's structure enables to incorporate individual and collective risks combined with proper equity, fairness and safety constraints, which leads to welfare generating policies. Contrary to traditional linear programming [2] and general equilibrium approaches [19], the model allows to deal with economies of scales, time dynamics and increasing returns. This phenomenon is typical for practical problems of production and resource allocation, however the discussion of these topics is beyond the scope of the paper. In contrast to general equilibrium and standard growth theory, the proposed risk-adjusted approach permits to deal with issues involving externalities, inherent uncertainties,

non-monetary values such as environmental degradation, non-marketable risks of high consequences, social heterogeneities regarding various representative agents.

Allocation of production facilities have to reasonably confirm to the distribution of current and future consumers including evaluation of the option make-vs.-buy typically addressed in spatial production planning models [15, 18, 20]. This implies the analysis of main production and demand driving forces such as population growth, urbanizations, energy provision, infrastructure, markets and market access. The discussed model can easily address regional “behavioral” aspects of production planning if these are determined by criteria other than pure cost-benefit or risks analysis. For example, rebalancing production allocation procedure in Sect. 10.3.3 allows to account for heterogeneous cultural traditions, complex interactions of behavioral, socio-economic, cultural and technological factors [7, 26], or specific fairness and equity considerations [23].

Within a project on “Policy Decision Support for Sustainable Adaptation of China’s Agriculture to Globalization” (CHINAGRO [17]), the model included specifics of China agricultural developments and has been applied for the spatial analysis of future livestock sector expansions. Using alternative economic and demographic projections [4, 16, 17, 21, 22], the model estimates per capita demand increases and consumption of major agricultural products, e.g., cereals, meat, milk, etc. Demand patterns differ between urban and rural areas, between geographical regions, and vary with income. Thus, with increasing incomes higher quality low-fat meat, e.g., poultry is preferred. In fact, evolution of consumption is modeled as a function of group-specific per capita income increases by applying income elasticities and distinguishing urban and rural consumers.

Agricultural supply is represented at county level, i.e., for about 2,430 spatial units. Smallholders and specialized livestock farms adjust the livestock herd structure and production in response to the demand increase and the changes of consumption patterns. The model distinguishes the following livestock types: poultry, pigs, dairy, cattle, buffaloes, yaks, sheep and goats, and other large animals (combining horses, donkeys, and camels). To examine the current situation and the production intensification trends, modeling of livestock production considers three management systems: traditional, specialized/industrial, and grazing.

In the environmental module, the environmental loads caused by intensive crop and livestock production are evaluated against admissible environmental and health thresholds (which can be proposed by stakeholders and environmental experts). Indicators used for measuring environmental impacts and human health risk are: the density of livestock, nutrients from manure and chemical fertilizers in excess of a location’s nutrient uptake by crop production, urbanization share, density of population, and others. Combinations of these and other factors (see Sect. 10.5) reflect different degrees of socio-economic and environmental risk exposures and can be used to guide sustainable production allocation. The model simulates different paths of demand increase, which induces respective location-specific production adjustments. In some locations, the environmental and health risk indicators may already exceed admissible thresholds, which signals that further production growth in these locations should not take place. This raises the question of how to adjust

the composition and allocation of livestock production facilities in response to increasing demand but without exacerbating environmental and health problems. The detailed description of the model and the allocation procedure [12] is rather lengthy. Therefore, in the following we provide only rather aggregate representation of their main constraints.

10.3.2 A Simplified Production Model

When planning livestock development, the objective is to allocate the foreseeable increases of demand for livestock products among the locations and the main production systems in the best possible way while accounting for various risks. In the following model the risks are treated as constraints restricting production expansion. In Sect. 10.4 we introduce a stochastic model that allows to account for risks and uncertainties in a more explicit manner.

Denote the expected national demand increase (to be satisfied by supply increase) in livestock product i by d_i , $i = \overline{1, m}$. Let x_{ijl} be the unknown supply increase in product i at location j and by management system l . In its simplest form, the problem is to find x_{ijl} satisfying the following system of equations:

$$\sum_{l,j} x_{ijl} = d_i, \quad (10.4)$$

$$x_{ijl} \geq 0 \quad (10.5)$$

$$\sum_i x_{ijl} \leq b_{jl}, l = \overline{1, L}, j = \overline{1, n}, i = \overline{1, m}, \quad (10.6)$$

where b_{jl} are thresholds aggregating environmental and health risks and imposing limitations to expand production in system l and location j . Equation (10.6) restricts prevalence of specific production systems. For example, the dominance of industrial systems in a location inevitably leads to intensification of feeding operations, the need in recycling facilities, etc. For simplicity of presentation, the constraint (10.6) captures only production side. Apart from b_{jl} , there may be additional limits on x_{ijl} , $x_{ijl} \leq r_{ijl}$, which can be associated with legislation, for example, to restrict production within a production belt, or to exclude from urban or protected areas, etc. Thresholds b_{jl} and r_{ijl} may either indicate that livestock in excess of these values is strictly prohibited or it incurs penalties such as taxes or premiums, for mitigation of the risks, say, livestock disease outbreaks or environmental pollution. Equations (10.4)–(10.6) belong to the type of transportation problems. However, there may be more general constraints of type $\sum_{ij} a_{ijl} x_{ijl} \leq d_i$ as in Sect. 10.2, $0 \leq a_{ijl} \leq 1$, which require extensions of the proposed approach.

In general, there exist infinitely many solutions of (10.4)–(10.6). The aim is to derive a solution that ensures appropriate balance between the efficiency and the risks. We can distinguish two sources of uncertainties generating potential risks: behavioral or endogenous uncertainties associated with allocation of new production capacities and exogenous uncertainties related to parameters of the model. In this section we consider only the first type of uncertainties. Section 10.4 addresses the second type of constraints.

The information on the current production facilities, threshold values b_{jl} , r_{ijl} , and costs are used to derive a prior probability q_{ijl} reflecting our belief that a unit of demand d_i should be allocated to management system l in location j . The use of priors is consistent with spatial economic theory (see discussion, e.g., in [7, 20, 26]). The likelihood q_{ijl} can be inversely proportional to production costs and inherent risks r_{ijl} [6, 7]. In Sect. 10.3.3 we show how it is used in a rebalancing procedure to determine the solution of (10.4)–(10.6) relying on behavioral, in a sense, risk-averse and cost-minimizing principles defined by this prior as in (10.10).

10.3.3 A Rebalancing Production–Allocation Algorithm

For simplicity of exposition, let us renumerate all pairs (l, j) , $l = \overline{1, L}$, $j = \overline{1, n}$ by $k = \overline{1, K}$. In this new notation, the problem is formulated as finding y_{ik} satisfying constraints:

$$\sum_k y_{ik} = d_i \tag{10.7}$$

$$y_{ik} \geq 0, \tag{10.8}$$

$$\sum_k y_{ik} = b_k, i = \overline{1, m}, k = \overline{1, K} \tag{10.9}$$

consistent with a prior q_{ik} belief that a unit of demand for product i should be supplied by activity k . For instance, it is reasonable to allocate more livestock to locations with a larger demand increase, higher productivity, or better feed access. Assume that this preference structure is expressed in prior q_{ik} , $\sum_k q_{ik} = 1$ for all i . In this case, the initial amount of production i allocated to k can be derived as $q_{ik}d_i$. But this may lead to violation of constraints (10.9). Sequential rebalancing [11] proceeds as follows. Relying on prior probability q_{ik} , the *expected* initial allocation of d_i to k is $y_{ik}^0 = q_{ik}d_i$, $i = \overline{1, m}$. However, this allocation may not satisfy constraint $\sum_i y_{ik}^0 \leq b_k$, $j = \overline{1, n}$. Derive the relative imbalances $\beta_k^0 = b_k / \sum_i y_{ik}^0$ and update $z_{ik}^0 = y_{ik}^0 \beta_k^0$, $i = \overline{1, m}$. Now the constraint $\sum_i y_{ik} \leq b_k$ is satisfied, $k = 1, 2, \dots$, but the estimate z_{ik}^0 may cause imbalance for (10.7), i.e., $\sum_k z_{ik}^0 \neq d_i$.

Continue with calculating $\alpha_i^0 = d_i / \sum_k z_{ik}^0$, $i = \overline{1, m}$, and updating $y_{ik}^1 = z_{ik}^0 \alpha_i^0$, an so on. The estimate y_{ik}^s can be represented as

$$q_{ik}^s = (q_{ik} \beta_k^{s-1}) / (\sum_j q_{ik} \beta_k^{s-1}),$$

where $i = \overline{1, m}$, $k = 1, 2, \dots$

Assume $y^s = y_{ik}^s$ has been calculated. Find $\beta_k^s = \bar{b}_k / \sum_i y_{ik}^s$ and $q_{ik}^{s+1} = (q_{ik} \beta_j^s / \sum_i q_{ik} \beta_j^s)$, $i = \overline{1, m}$, $k = 1, 2, \dots$, and so on.

In this form the procedure can be viewed as a redistribution of required supply d_i among producers $k = 1, 2, \dots$ by applying sequentially adjusted q_{ik}^{s+1} , e.g., by using a Bayesian type of rule for updating the prior distribution:

$$q_{ik}^{s+1} = q_{ik} \beta_k^s / \sum_i q_{ik} \beta_k^s,$$

were $q_{ik}^0 = q_{ik}$.

The update is done on an observation of imbalances of basic constraints rather than observations of random variables. A rebalancing procedure, similar to the one described above for Hitchcock–Koopmans transportation constraints (10.7)–(10.9), was proposed by G.V. Sheleikovskii (see a proof and references in [3]) for estimation of passenger flows between regions. A proof of its convergence to the optimal solution maximizing the cross-entropy function

$$\sum_{i,k} y_{ik} \ln \frac{y_{ik}}{q_{ik}} \tag{10.10}$$

is given in [11] for rather general types of constraints. It should be noted that in our model we use equality constraints (10.9). The general inequality constraints are reduced to this model by introduction of a fictitious demand constraint.

10.4 Stochastic Production Allocation Model

The approach presented in Sect. 10.3 evaluates the increase of livestock production relying on individual behavioral principles set by priors. There, the risks are characterized in a simplified deterministic way by imposing certain standards as additional “safety” constraints. In general, these constraints may depend on some scenarios of potential future shocks. The behavioral uncertainty in Sect. 10.3 can also be treated in a stochastic manner as allocation of random demand $d_i(\omega)$ among points $k = \overline{1, K}$ with respect to the prior q_{ik} , which is a topic of a separate paper.

Let us consider now a more general multi-producer model in a stochastic environment analogous to the Example of Sect. 10.2. We may assume that there is a

coordinating agency. The goal of this agency is to maximize the overall performance of the production chain with large and small units to stabilize the aggregate production and increase the facility values. Suppose that the agency has to determine levels of livestock product i in locations k in order to meet stochastic demand $d_i(\omega)$, where $\omega = (\omega_1, \omega_2, \dots)$ is a vector of all contingencies affecting demand and production. Naturally to assume that the decision on production expansion has to be made before the information on contingencies arrives. In this case, the total ex-ante production may not exactly correspond to the real demand, i.e., we may face both oversupplies and shortfalls. In other words, the amount of production $y_{ik}, k = \overline{1, K}$, which is planned ex-ante to satisfy the demand $d_i(\omega)$, $y_i(\omega) = \sum a_{ik}(\omega)y_{ik}$ may underestimate ($y_i(\omega) < d_i(\omega)$) or overestimate ($y_i(\omega) > d_i(\omega)$) the real demand $d_i(\omega)$ under revealed contingencies and the safety constraints imposed by strict thresholds b_k in (10.9). The constraint (10.9) necessitates, in general, additional supply of ex-ante production $z_i \geq 0$ from external sources (say, through international trade). It may also require the ex-post redistribution of the production from internal producers, $k = \overline{1, K}$, to eliminate arising shortfalls and oversupplies in locations. For now, let us ignore these ex-post redistributive aspects assuming that the most significant impacts are associated with ex-ante decisions y_{ik} and z_i . In fact, the presented further model can be easily extended to represent the ex-post adjustments of decisions y_{ik}, z_i , as well as temporal aspects of production planning.

Let c_{ik} be the unit production cost. In more general model formulation, c_{ik} may also include the unit transportation cost for satisfying location-specific demand. Then the model of production planning among the facilities can be formulated as the minimization of the total cost function:

$$f(y, z) = \sum_{i,k} c_{ik}y_{ik} + \sum_{i=1}^m e_i z_i,$$

subject to constraints (10.8), (10.9), and the following additional safety constraints

$$P\left[\sum_{k=1}^K a_{ik}(\omega)y_{ik} + z_i \geq d_i(\omega)\right] \geq p_i, z_i \geq 0, i = \overline{1, m}, \tag{10.11}$$

where $e_i > 0, i = \overline{1, m}$, denotes the unit import cost. A safety level $p_i, 0 < p_i < 1$, defines (ensures) the stability of the supply-demand relations for all possible scenarios (contingencies) ω . The introduction of constraints of type (10.11) is a standard approach for characterizing stability in case of the insurance business, operations of nuclear power plants and other risky activities especially when involving catastrophic risks [9]. Safety constraints of type (10.11) are usually used in cases where impacts of random interruptions can not be easily evaluated. In this case, the value p_i is selected such that an expected shortfall occurs only, say, once in 100 month, i.e., $p_i = 0.01$.

The main methodological challenge is concerned with the lack of convexity of constraints (10.11). Yet, the remarkable fact is that the model defined by

(10.8)–(10.11) can be effectively solved by linear programming methods due to the following equivalent convex form of this model. Let us consider the minimization of the expectation function

$$F(y, z) = f(y, z) + \sum_{i=1}^m \alpha_i E \max\{0, d_i(\omega) - \sum_{k=1}^K a_{ik}(\omega)y_{ik} - z_i\}, \quad (10.12)$$

subject to constraints (10.8), (10.9), and $z_i \geq 0, i = \overline{1, m}$. The minimization of function $F(y, z)$ is a rather specific case of stochastic minimax models analyzed (both optimality conditions and solution procedures) in [8]. In particular, if $F(y, z)$ has continuous derivatives with respect to z_i , e.g., the probability distribution function of ω has continuous density function, then

$$\frac{\partial F}{\partial z_i} = e_i - \alpha_i E I(d_i(\omega) - \sum_{k=1}^K a_{ik}(\omega)y_{ik} - z_i \geq 0)$$

where $I(\xi \geq 0)$ is the indicator function: $I(\xi \geq 0) = 1$, if $\xi \geq 0$, and $I(\xi \geq 0) = 0$ otherwise. Therefore, we can rewrite $\frac{\partial F}{\partial z_i}$ as

$$\frac{\partial F}{\partial z_i} = e_i - \alpha_i P[d_i(\omega) - \sum_{k=1}^K a_{ik}(\omega)y_{ik} - z_i \geq 0], \quad (10.13)$$

which allows to establish connections between the original model defined by (10.8)–(10.11) and the minimization of convex function $F(y, z)$ defined by (10.12).

Assume (y^*, z^*) minimizes $F(y, z)$ subject to constraints (10.8), (10.9), and $z_i \geq 0, i = \overline{1, m}$. Assume also that $e_i < \alpha_i, i = \overline{1, m}$. Then from (10.13) it follows that for all i with positive components $z_i^* > 0$, i.e., when $\frac{\partial F}{\partial z_i} = 0$, the optimal solution (y^*, z^*) satisfies the following safety constraints

$$P[d_i(\omega) - \sum_{k=1}^K a_{ik}(\omega)y_{ik} - z_i \geq 0] = e_i/\alpha_i. \quad (10.14)$$

Moreover, for all i with $z_i^* = 0$, i.e., when $\frac{\partial F}{\partial z_i} \geq 0$, the optimal (y^*, z^*) satisfies the following safety constraint

$$P[d_i(\omega) - \sum_{k=1}^K a_{ik}(\omega)y_{ik} \geq 0] \leq e_i/\alpha_i. \quad (10.15)$$

If we choose α_i as $e_i/\alpha_i = 1 - p_i$, i.e., $\alpha_i = e_i/(1 - p_i)$, then (10.14)–(10.15) become equivalent to the safety constraint (10.11) of the original model (10.8)–(10.11). In other words, the minimization of convex function $F(y, z)$ defined by

(10.12) subject to (10.8), (10.9), and $z_i \geq 0, i = \overline{1, m}$, yields the optimal solution of the original model (10.8)–(10.11). Efficient computational procedures for solving stochastic minimax problems with objective functions defined as in (10.12) can be found in [8, 25]. In particular, the paper [25] discussed the applicability of linear programming methods in cases where the original model defined by a general probability distributions of ω can be sufficiently approximated by models with discrete probability distributions. This paper establishes also important connections between the minimization of (10.12)-type functions and Conditional-Value-at-Risk risk measure.

The minimization of function (10.12) can also be solved by a stochastic quasi-gradient method [8]. In applying this method to minimization of (10.12), the differentiability of $F(y)$ and any assumption on probability distribution of ω is not required. Also, the probability distribution of ω may only be given implicitly. For instance, only observations of random $d_i(\omega)$ and $a_{ik}(\omega)$ may be available or only a Monte Carlo procedure (pseudo-sampling simulation model such as described in Sect. 10.3.1) is used to simulate supply and demand. In the following section we illustrate some applications by using only the rebalancing algorithm described in Sect. 10.3.3; elaboration of the outlined stochastic allocation algorithm is a topic for future implementation.

10.5 Numerical Experiments

The model in Sect. 10.3 is used in the analysis of current and plausible future livestock production allocation and intensification in China. Namely, in each time period the simulation model generates levels and geographic distribution of demand for livestock products coherent with urbanization processes [22], demographic change [4] and expected growth of incomes [16, 17]. Production allocation and intensification levels are projected from the base year data for the main livestock types (pigs, poultry, sheep, goat, cattle) and management systems (grazing, industrial/specialized, traditional) at the level of counties (about 2,500 administrative units). For production allocation, we used the sequential rebalancing procedure described in Sect. 10.3.3. Two scenarios of future production allocation corresponding to different priors $q_{ik}, i = \overline{1, m}, k = \overline{1, K}$, are compared: (1) an intensification scenario, when production is allocated proportionally to the geographical patterns of demand increases, and (2) a risk-adjusted scenario that combines the preference structure as defined by the geographical distribution of demand with indicators of environmental pressure.

Intensification scenario. Currently, common practice is to allocate intensive livestock production in areas with good access to consumers, close to high demand and high population density [1]. In many practical problems of large dimensionality, to describe the “profitability” of a location it has been standard practice to use an ad hoc but reasonable measure referred to as market access function. The typical market access function measures the potential of location k as a weighted sum of purchasing power of all other locations in some vicinity of the given k . The weights

are defined either as a function of distance or as a function of other factors, say, costs or losses. In these studies, each county is characterized by its market access calculated as a weighted sum of demand for product i in nearby counties within some vicinity. Values $\tilde{\Delta}_{ik}$, determine a profit-based prior q_{ik} , $k = \overline{1}, \overline{K}$ for allocation of demand increase among production units in locations as it is described in Sect. 10.3.3.

Risk-adjusted scenario. The objective of this scenario is to care for the balance between profitability of the agricultural production, rural welfare, and the respect of nature and the environment. Challenges of spatially-explicit planning for sustainability are related to the choice of adequate location-specific indicators to guide rural development within defined socio-economic and environmental objectives. While information on economic and livelihood conditions at location may be available from statistics and census data, estimation of agricultural pollution and health risks (from statistics and census data, estimation of agricultural pollution and health risks (from example, related to livestock diseases) is a more challenging task.

The agricultural pollution falls into the category of non-point source pollution, which is geographically disperse, and the likelihood of disease occurrences is determined by a combination of factors. Measurements of the pollution level, health risks, and related impacts or losses are hardly possible as they depend on multiple highly uncertain socio-economic and environmental factors: weather patterns, population density, level of development, agricultural inputs and intensification levels, etc. In many practical situations when the target variable is impossible or impractical to measure, it is possible to use context-specific proxies or even a set of proxies that can considerably well represent the state of the non-measurable variable (see, e.g., [13]).

For planning sustainable agricultural developments, the profit-driven prior of the “intensification scenario” is adjusted with such variables as nutrients in excess of crop uptakes, density of livestock biomass, etc., are used to characterize environmental risks. Health norms and associated health risks are introduced by a combination of urbanization share (share of urban population in total population) and availability of non-residential area suitable for further production expansion in each location. In general, allocation prior is defined by a compound probability distribution function of relevant variables.

The intensification scenario implicitly minimizes the transportation costs as the production concentrates in the proximity of large markets in urban areas with high demand. In the alternative scenario, which is a compromise between the demand driven production allocation and the considerations of health and environmental risks, the production is shifted to more distant locations characterized by availability of cultivated land, lower livestock and population density, which increases transportation. However, the measure of goodness for the scenarios accounts not only for the transportation cost but also includes environmental and health risk proxies. Thus, the two scenarios are compared with respect to number of people in China’s regions exposed to different categories of environmental risks.

Environmental risks are measured in terms of environmental pressure in relation to the coincidence of three factors: density of confined livestock, human population density, and availability of cultivated land. For this purpose some 2,434 counties

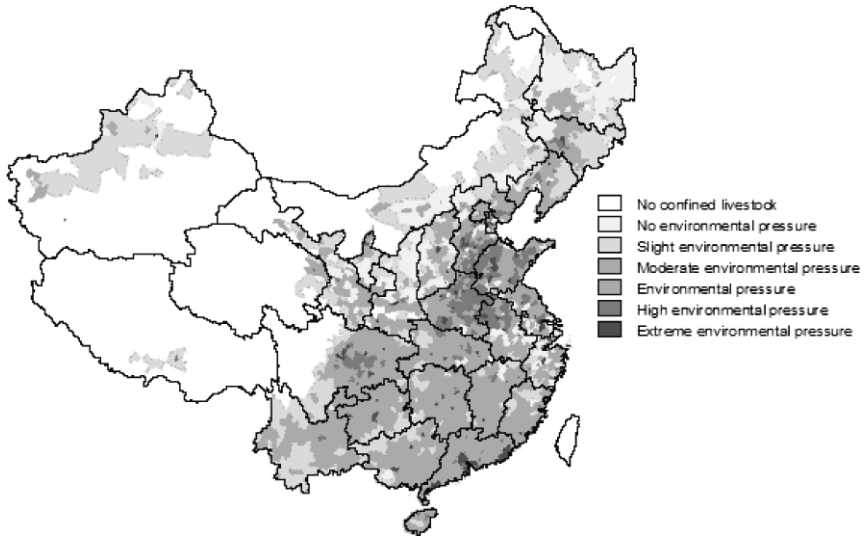


Fig. 10.1 Environmental pressure from confined livestock production, 2000

were classified as follows into seven categories, namely: (a) *No confined livestock*, i.e., counties in scarcely populated areas (desert or mountain/plateau) and with very little confined livestock; (b) *No environmental pressure*, i.e., counties with substantial crop production but with little confined livestock; (c) *Slight environmental pressure* counties with low environmental pressure from confined livestock production; (d) *Moderate environmental pressure*, i.e., counties with moderate environmental pressure from confined livestock production; (e) *Environmental pressure*, i.e., counties with substantial urbanization and environmental pressure from confined livestock production; (f) *High Environmental pressure*, i.e., counties with substantial urbanization and high environmental pressure from livestock production, and (g) *Extreme environmental pressure* i.e., counties with high degree of urbanization coinciding with high environmental pressure from confined livestock production. Figure 10.1 presents the above classification of environmental pressure for the year 2000. Figure 10.1 indicates that currently (i.e., year 2000) hot-spots of environmental pressure are located mainly in provinces covering the North China Plain, the Sichuan basin, and several locations along the coast of South China. Locations of livestock production concentrate around or in the vicinity of areas where the livestock demand grows fast, e.g., highly populated and urban areas.

Figure 10.2 presents diagrams of the distribution of current population against the mapped classes of severity of environmental pressures from livestock. The left diagram shows absolute numbers, i.e., million people per class and region. The diagram on the right gives shares of population within each region falling into respective classes. For year 2000, the estimates suggest that about 20% of China’s population lives in counties characterized as having high or extreme severity of environmental pressure from intensive livestock production. In the “intensification” scenario, by

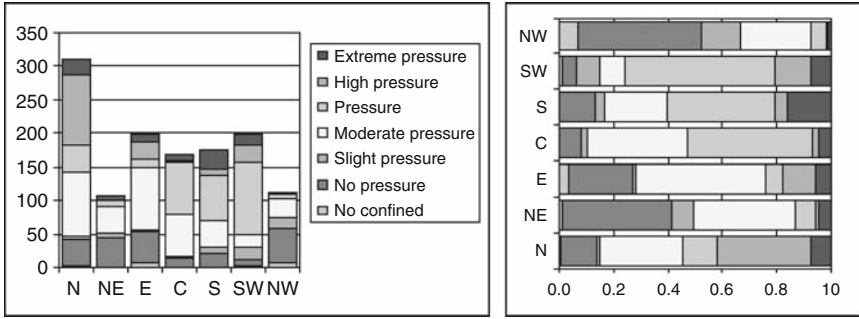


Fig. 10.2 Absolute (million people) and relative (share of total population) distribution of population according to classes of severity of environmental pressure from livestock, 2000. The label on the horizontal axis indicate China regions: N, NE, E, C, S, SW, NW stand for North, North-East, East, Center, South, South-West, North-West, respectively

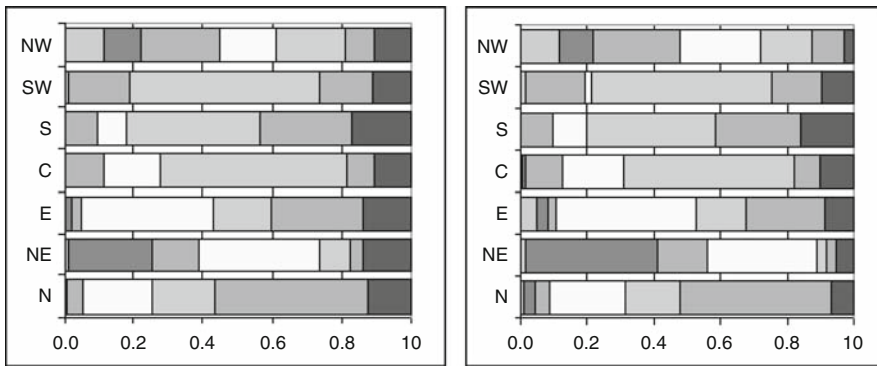


Fig. 10.3 Relative (share of total population) distribution of population according to classes of severity of environmental pressure from livestock, 2030: **a** “intensification” scenario, **b** risk-adjusted scenario

2030 this population share increases to 36% (Fig. 10.3a), i.e., from one-fifth in 2000 to about one-third in 2030. Looking only at the highest pressure class, the South region appears to have the largest number of people and the highest population share in such unfavorable environmental pressure, about 38 million or 22% of population in 2000 increasing to nearly 45 million or 17% in 2030. The region with the highest occurrence of people (both absolute and relative) in the two highest pressure classes is the North region, with more than 40% of the population. In 2030, the estimated share becomes 57%, followed by the South region with 27% population in two highest pressure classes in 2000 and with 44% in 2030. Looking at the second allocation scenario, the positive changes are quite visible (see Fig. 10.3b). The estimate of people living in highest pressure class for South region changes from 45 to 42 million. Percentage of population in the two highest pressure classes for the same region varies between scenarios as 43 for the bad and 41 for the good. For

North region, 56.5% of total population will live in two highest pressure classes in 2030 in bad scenario and only 52.3 – in good. In North East, the highest pressure classes change percentage from about 18 to less than 10. The intensification scenario (1) implicitly minimizes the transportation costs as production concentrates in the vicinity of urban areas with high demand. In the alternative scenario (2), the production is shifted to more distant locations characterized by availability of cultivated land, lower livestock and population density, but at the expense of additional transportation. Environmental sustainability aspects of the two scenarios were compared with respect to the share of people in China's regions exposed to different severity classes of environmental risks. Environmental risks are measured in terms of environmental pressure in relation to the coincidence of three factors: density of confined livestock, human population density, and availability of cultivated land. For year 2000, the estimates suggest that about 20% of China's population lives in counties characterized as having high or extreme severity of environmental pressure from intensive livestock production. In the "intensification" scenario, by 2030 this population share increases to 37%, while in the second, environmentally friendly scenario, it stays below 30%. To finally compare the two scenarios, it is necessary to "normalize" gains due to improved life conditions with expenses of additional transportation.

10.6 Conclusions

This paper addresses some important aspects of agricultural production planning under risks, uncertainties and incomplete information. When planning agricultural developments, the objective is to allocate the foreseeable increases of demand in the best possible way while accounting for various risks associated with production and suitability criteria for profitability, transport, health and environmental impacts. Models for production allocation under risks and uncertainties may have considerable implications. In particular, the allocation of livestock production away from urban peripheries where pressure is highest to regions where feed grains are in abundance could decrease the income gaps between the regions. Similarly, establishment of agricultural pollution regulations, e.g., taxation, at locations with high environmental loads may change the balance of agricultural market attracting imports from abroad.

In Sect. 10.3, the production allocation procedure is proposed for situations when the available information is given in the form of aggregate values without providing necessary local perspectives. Therefore, the main issue is to downscale these values to the local levels consistently with location specific behavioral principles based on some priors. Yet, many practical situations may require more rigorous probabilistic treatment of priors and safety constraints. Section 10.4 proposes an allocation mechanism with more general treatment of uncertainties and risks based on principles of stochastic optimization. This is a promising approach for a coordinating agency aiming to improve the overall performance of the production chain.

By diversifying large and small units the agency may stabilize the aggregate production and increase “utility” of individual facilities. The application of this allocation procedure is a topic for future research.

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Chapter 11

Dealing with Uncertainty in GHG Inventories: How to Go About It?

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Abstract The assessment of greenhouse gases emitted to and removed from the atmosphere is high on both political and scientific agendas. Under the United Nations Framework Convention on Climate Change, Parties to the Convention publish annual or periodic national inventories of greenhouse gas emissions and removals. Policymakers use these inventories to develop strategies and policies for emission reductions and to track the progress of these policies. However, greenhouse gas inventories (whether at the global, national, corporate, or other level) contain uncertainty for a variety of reasons, and these uncertainties have important scientific and policy implications. For scientific, political, and economic reasons it is important to deal with the uncertainty of emissions estimates proactively. Proper treatment of uncertainty affects everything from our understanding of the physical system to the economics of mitigation strategies and the politics of mitigation agreements. A comprehensive and consistent understanding of, and a framework for dealing with, the uncertainty of emissions estimates should have a large impact on the functioning and effectiveness of the Kyoto Protocol and its successor. This chapter attempts to

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pull together relevant fragments of knowledge, allowing us to get a better picture of how to go about dealing with the uncertainty in greenhouse gas inventories.

11.1 Introduction

The assessment of greenhouse gases (GHGs) emitted to and removed from the atmosphere is high on both political and scientific agendas. Under the United Nations Framework Convention on Climate Change (UNFCCC), Parties to the Convention (so-called Annex I countries) have published annual or periodic national inventories of GHG emissions and removals since the mid 1990s. Policymakers use these inventories to develop strategies and policies for emission reductions and to track the progress of these policies. Where formal commitments to limit emissions exist, regulatory agencies and corporations rely on inventories to establish compliance records. Scientists, businesses, the public, and other interest groups use inventories to better understand the sources and trends in emissions. Table 11.1 provides general background information on the six GHGs, or groups of gases, considered under the Kyoto Protocol and their global emissions as reported by the Intergovernmental Panel on Climate Change (IPCC) in its assessment reports for the late 1990s and beyond.

GHG inventories contain uncertainty for a variety of reasons – for example, the lack of availability of sufficient and appropriate data and the techniques to process them. Uncertainty has important scientific and policy implications. However, until recently, relatively little attention has been devoted to how uncertainty in emissions estimates is dealt with and how it might be reduced. Now this situation is changing, with uncertainty analysis increasingly being recognized as an important tool for improving national, sectoral, and corporate inventories of GHG emissions and removals [5] (see also [6] and [7]).

At present, Parties to the UNFCCC are encouraged, but not obliged, to include with their periodic reports of in-country GHG emissions and removals, estimates of the uncertainty associated with these emissions and removals; consistent with the Intergovernmental Panel on Climate Change's (IPCC) good practice guidance reports [8,9]. Inventory uncertainty is monitored, but not regulated, under the Kyoto Protocol [5].

We argue that it makes a big difference in the framing of policies whether or not uncertainty is considered: reactively, because there is a need to do so; or proactively, because difficulties are anticipated. We follow [7, p. 2–3] (see also [5]) who clearly state that uncertainty estimates are not intended to dispute the validity of national GHG inventory figures. Although the uncertainty of emissions estimates underscores the lack of accuracy that characterizes many source and sink categories, its consideration can help to establish a more robust foundation on which to base policy.

According to the IPCC good practice guidance reports (notably, [8, p. 6.5]), uncertainty analysis is intended to help “improve the accuracy of inventories in the future and guide decisions on methodological choice.” Uncertainty analyses function as indicators of opportunities for improvement in data measurement, data

Table 11.1 The six GHGs, or groups of gases, considered under the Kyoto Protocol to the UNFCCC [1, Annex A] and their global emissions as reported by the IPCC in its Third and Fourth Assessment Reports for the late 1990s and beyond. The GWP (last column) describes the global warming potential for a given GHG. It allows expressing the emissions of a non-CO₂ GHG in terms of CO₂-equivalent, which is the amount of CO₂ that causes the same global warming when measured over a specified timescale (generally, 100 years). The relative uncertainty ranges within which Annex I countries generally report the emissions of these GHGs are specified in Table 11.3

Kyoto gas	Global emissions		GWP ^a [3, Table TS.2]
	[2, Table 4.1], [4, Tables 7.1, 7.6, 7.7]		
	Anthropogenic late 1990s ^b	Natural late 1990s ^b	
Carbon dioxide (CO ₂)	2000–2005: 7.2 ± 0.3 Pg C 26.4 Pg CO ₂ -eq.	2000–2005: –3.1 ± 0.8 Pg C ^c 11.4 Pg CO ₂ -eq.	1
	1996–2001: 428 Tg ^d 9.0 Pg CO ₂ -eq.	1996–2001: 168 Tg ^e 3.5 Pg CO ₂ -eq.	21 (25)
Methane (CH ₄)	1990s: 6.7 Tg N ^f 6.5 Pg CO ₂ -eq.	1990s: 11.0 Tg N ^g 10.7 Pg CO ₂ -eq.	310 (298)
	See below	None	See below
Perfluorocarbons (PFCs)	See below	Negligible	See below
Sulfur hexafluoride (SF ₆)	~6 Gg 0.14 Pg CO ₂ -eq.	Negligible	23,900 (22,800)
<i>Important HFCs and PFCs:</i>			
HFC-23 (CHF ₃)	~7 Gg 0.08 Pg CO ₂ -eq.	None	11,700 (14,800)
HFC-134a(CH ₂ FCF ₃)	~25 Gg 0.03 Pg CO ₂ -eq.	None	1,300 (1,430)
HFC-152a (CH ₃ CHF ₂)	~4 Gg 0.56 Tg CO ₂ -eq.	None	140 (124)
PFC-14 (CF ₄)	~15 Gg 0.10 Pg CO ₂ -eq.	Negligible	6,500 (7,390)
PFC-116 (C ₂ F ₆)	~2 Gg 0.02 Pg CO ₂ -eq.	None	9,200 (12,200)

^a The net global warming potential (GWP) refers to a time horizon of 100 years. The GWPs stem from the IPCC Second Assessment Report (Climate Change 1995) as these are used for reporting under the UNFCCC. The most recent GWP updates for the IPCC Fourth Assessment Report (Climate Change 2007) are reported in addition (in parentheses)

^b If not indicated otherwise

^c Uptake: net atmosphere-to-land flux; and atmosphere-to-ocean flux

^d Emissions: coal mining; gas, oil and industry; ruminants; rice agriculture; and biomass burning

^e Emissions: wetlands and termites

^f Emissions: fossil fuel combustion and industrial processes; agriculture; biomass and biofuel burning; human excreta; rivers, estuaries and coastal zones; and atmospheric deposition

^g Emissions: soils under natural vegetation; oceans; and atmospheric chemistry

collection, and calculation methodology. Only by identifying elements of high uncertainty can methodological changes be introduced to address them. Currently, most countries that perform uncertainty analyses do so for the express purpose of improving their future estimates; and the rationale is generally the same at the corporate and other levels. Estimating uncertainty helps to prioritize resources and to take precautions against undesirable consequences. Depending on the intended purpose of an inventory, however, this may not be the extent of the utility of uncertainty analysis. Another rationale for performing uncertainty analysis is to provide a policy tool, a means to adjust inventories or analyze and compare emission changes in order to determine compliance or the value of a transaction. While some experts find the quality of uncertainty data associated with national inventories insufficient to use for these purposes, others offer justification for conducting uncertainty analyses to inform and enforce policy decisions. Some experts suggest revising the system of accounting on which current reduction schemes are based, while others seek to incorporate uncertainty measurements into emission and emission change analysis procedures. The latter could offer policy makers enhanced knowledge and additional insight on which to base GHG emission reduction measures.

We follow the proactive track in dealing with uncertainty. In Sect. 11.2 we look into the question of why uncertainty matters in general. Sections 11.3 and 11.4 elaborate on Sect. 11.2. In Sect. 11.3 we provide an overview of the state-of-the-art of analyzing emission changes in consideration of uncertainty. We envision this analysis taking place in accordance with, not independent of, a dual-constrained (bottom-up/top-down) verification framework in Sect. 11.4. We summarize our findings in Sect. 11.5.

11.2 Does Uncertainty Matter?

Reference [5] (see also [7]) offers a number of reasons why the consideration of uncertainty in GHG inventories is important:

- Understanding the basic science of GHG sources and sinks requires an understanding of the uncertainty in their estimates.
- Schemes to reduce human-induced global climate change rely on confidence that inventories of GHG emissions allow the accurate and transparent assessment of emissions and emission changes.
- Uncertainty is higher for some aspects of a GHG inventory than for others. For example, past experience shows that, in general, methods used to estimate nitrous oxide (N₂O) emissions are more uncertain than those for methane (CH₄) and much more uncertain than those for carbon dioxide (CO₂). Whether in multi-gas, cross-sectoral, international comparisons, trading systems, or in compliance mechanisms, approaches to uncertainty analysis need to be robust and standardized across sectors and gases, as well as among countries.

Uncertainty analysis helps to understand uncertainties: better science helps to reduce them. Better science needs support, encouragement and investment. Full carbon accounting (FCA) – or full accounting of emissions and removals, including all

GHGs – in national GHG inventories is important for advancing the science. FCA is a prerequisite for reducing uncertainties in our understanding of the global climate system. From a policy viewpoint, FCA could be encouraged by including it in reporting emissions, but it might be separated from targets for reducing emissions. Future climate agreements will become more robust if there is explicit accounting for the uncertainties associated with emission estimates. Hence, understanding uncertainty matters in many ways.

11.3 State of the Art of Analyzing Uncertain Emission Changes

In this section we elaborate on Sect. 11.2 by looking into the state-of-the-art of analyzing changes in emissions and removals of GHGs in consideration of uncertainty. From a physical (measurability) point of view, the uncertainty surrounding emission changes becomes more important in relative terms the smaller are the changes in the emissions, that is, the smaller the dynamics that they exhibit. Two options exist to avoid situations of great uncertainty vs. small change: (1) allowing more time so that greater emission changes can materialize; and (2) increasing measurability, e.g., by focusing on GHG emissions that can be grasped with “sufficient” certainty so that their changes are still “significant” in spite of the uncertainty. (Alternatively, emissions that do not possess these characteristics should be treated differently, e.g., separately from single-point emission targets (see above), or only in connection with targets that are defined as emission intervals or corridors.) Given that renegotiating the commitment times under the Kyoto Protocol cannot happen, Option 1 is not considered further. Option 2 requires the application of techniques that allow analyzing emission changes quantitatively (i.e., on an intra-technique basis) and qualitatively (i.e., on an inter-technique basis). Any of these techniques can be applied to GHG emissions individually, that is, they allow a detailed and thorough comparison of agreed or realized changes in emissions.

While pursuing the analysis of uncertain emission changes (also termed emission signals), we typically refer to the country scale, the principal reporting unit for reporting GHG emissions and removals under the Kyoto Protocol, but we could also refer to any other spatio-thematic scale. Our main motivation for studying the uncertainty of country-scale emissions estimates is the still unresolved issue of compliance (see also [10]). For most countries the emission changes agreed on under the Kyoto Protocol are of the same order of magnitude as, or smaller than, the uncertainty that underlies their combined CO₂ equivalent emissions estimates (compare the right column of Table 11.2 with the second column of Table 11.3).¹ Techniques are not in place to analyze uncertain emission signals from various points

¹ The issue of great uncertainty vs. small change also arises for small, intermediate, reduction targets. For instance, the EU discusses annual reduction steps in the context of an overall (EU-wide) GHG emission reduction of 20% by 2020 compared to 2005 [11, p. 7]. These steps follow a linear reduction path and are small (<2% per year; not compounded).

Table 11.2 Countries included in Annex B to the Kyoto Protocol (KP) and their emission limitation and reduction commitments (commitment period for all countries: 2008–2012; for the ISO Country Code for country abbreviations see below). The individual commitments have to be seen in context, i.e., vis-à-vis the uncertainty that underlies the reporting of emissions at the country scale (see Table 11.3). *Sources:* [14, Article 3.8, Annex B], [1, Decision 11/CP.4], [15], [16], [17, National Inventory Submissions], [18, Sect. 2.b]

Country group	Annex B country	Base year(s) for CO ₂ , CH ₄ , N ₂ O (for HFCs, PFCs, SF ₆)	KP commitment %
1a	See below ^a	1990 (1995)	
	See below ^b	1990 (1990)	
1b	RO	1989 (1989)	92
1c	BG	1988 (1995)	
1d	SI	1986 (1995)	
2	US ^c	1990 (1990)	93
3a	JP	1990 (1995)	
	CA	1990 (1990)	94
3b	PL	1988 (1995)	
3c	HU	1985–1987 (1995)	
4	HR	1990 (1995)	95
5a	RU	1990 (1995)	100
5b	NZ, UA	1990 (1990)	
6	NO	1990 (1990)	101
7	AU	1990 (1990)	108
8	IS	1990 (1990)	110

^aCountry Group 1a: BE, CZ, DE, DK, EC (= EU-15; the EU-27 does not have a common Kyoto target), EE, ES, FI, GR, IE, LT, LU, LV, MC, NL, PT, SE, UK. Member States of the EU-27 but without individual Kyoto targets: CY, ML. Listed in the Convention's Annex I but not included in the Protocol's Annex B: BY and TR (BY and TR were not Parties to the Convention when the Protocol was adopted). BY requested becoming an Annex B country by amendment to the Kyoto Protocol at CMP 2 in 2006. (CMP = Conference of the Parties serving as the Meeting of the Parties to the Kyoto Protocol.) BY's base years and emission reduction commitments are 1990 (1995) and 92%, respectively

^bCountry Group 1a: AT, CH, FR, IT, LI, SK

^cCountry Group 2: The US has indicated its intention not to ratify the Kyoto Protocol. The US reports all its emissions with reference to 1990. However, information on 1990 in its national inventory submissions does not reflect or prejudice any decision that may be taken in relation to the use of 1995 as base year for HFCs, PFCs and SF₆ in accordance with Article 3.8 of the Kyoto Protocol

Abbreviations: AT Austria, AU Australia, BE Belgium, BG Bulgaria, BY Belarus, CA Canada, CH Switzerland, CY Cyprus, CZ Czech Republic, DE Germany, DK Denmark, EC European Community, EE Estonia, ES Spain, FI Finland, FR France, GR Greece, HR Croatia, HU Hungary, IE Ireland, IS Iceland, IT Italy, JP Japan, LI Liechtenstein, LT Lithuania, LU Luxembourg, LV Latvia, MT Malta, MC Monaco, NL Netherlands, NO Norway, NZ New Zealand, PL Poland, PT Portugal, RO Romania, RU Russian Federation, SE Sweden, SI Slovenia, SK Slovak Republic, TR Turkey, UA Ukraine, UK United Kingdom, US United States

Table 11.3 Emissions and/or removals of GHGs, or combinations of GHGs, classified according to their relative uncertainty ranges (reference: country scale). The bars of the arrows indicate the dominant uncertainty range for these emissions and removals, while the tops of the arrows point at the neighboring uncertainty ranges, which cannot be excluded but appear less frequently. LULUCF stands for the direct human-induced land-use, land-use change, and forestry activities stipulated by Articles 3.3 and 3.4 under the Kyoto Protocol [14]. The arrows are based on the total uncertainties that are reported for the Member States of the EU-25 [19] and the expertise available at IIASA's Forestry Program (cf. http://www.iiasa.ac.at/Research/FOR/unc_bottomup.html) and elsewhere (e.g., [20, Sects. 2.3.7, 2.4.1], [9, Sect. 5.2]). *Source:* [21, Table 1], modified

Class	Relative uncertainty (%) for 95% confidence interval	Classification of emissions and/or removals
1	0–5	↓ CO ₂ from fossil fuel (plus cement)
2	5–10	↕ All Kyoto GHGs
3	10–20	↕ Plus LULUCF
4	20–40	↓
5	>40	↓ CO ₂ net terrestrial (>80%)

of view, ranging from signal quality (defined adjustments, statistical significance, detectability, etc.) to the way uncertainty is addressed (trend uncertainty or total uncertainty). Any such technique, if implemented, could “make or break” compliance, especially in cases where countries claim fulfillment of their reduction commitments. As already mentioned above, inventory uncertainty is monitored, but not regulated, under the Protocol. It remains to be seen whether the current status of ignoring uncertainty and abstaining from specifying clear rules on how to go about it will survive in the long term (see Compliance under the Kyoto Protocol in [12]).

The analysis of emission signals in consideration of uncertainty can take three forms involving a multitude of techniques: (1) preparatory, (2) midway, and (3) retrospective signal analysis. Preparatory signal analysis allows generating useful knowledge that one would ideally wish to have at hand before negotiating international environmental agreements such as the Kyoto Protocol or its successors. For instance, it is important to know beforehand how great the uncertainties could be, depending on the desired level of confidence in the emission signal. What is the signal one wishes to detect and what is the risk one is willing to tolerate in meeting an agreed emission limitation or reduction commitment? To this end, it is generally assumed that (1) the emissions path between the base year and commitment year/period is a straight line (this is only a boundary condition, not a restriction); and (2) our knowledge of uncertainty in the commitment year/period will be as good as today's, in relative (qualitative) terms. Preparatory signal analysis allows factoring in the change in uncertainty, which can be due to learning and/or can result from structural changes in the emitters. However, researchers only begin to grasp these two determinants and to discriminate between them [22]. Handling the change in uncertainty is within reach but more data and research are needed. Being able to estimate the change in uncertainty is important in setting appropriate emission reduction targets, but one must not forget that preparatory signal analysis has not yet been applied in its simplest form to Kyoto commitments.

The state-of-the-art of preparatory signal analysis is well summarized by [23] (see also [21,24–26]), who compare six of the most widely discussed techniques.² In addition, preparatory signal analysis also allows monitoring the success of a country in reducing its emissions along a prescribed emissions target path between its base year and commitment year/period. This positive feature opens up a range of policy-relevant applications.³

Midway and retrospective signal analysis are less advanced than preparatory signal analysis. So far, midway signal analysis still focuses on emissions rather than on emissions changes. Midway signal analysis is an attempt to assess information on an emissions path at some point in time between base year and commitment. It considers a signal's path realized so far vis-à-vis a possible path toward the agreed emission limitation or reduction commitment. In this process, the dynamical moments (velocity, acceleration, etc.) of the historical and envisioned paths are compared, and this indicates (first-order control) whether or not it is likely to achieve the emission commitment. Midway signal analysis generally incorporates information from emissions prior to the base year to determine the signal's dynamical moments more accurately. The techniques explored so far to grasp the dynamics of, mostly, fossil-fuel CO₂ emissions encompass: polynomial regressions [29]; integral transforms [30]; and smoothing splines, parametric modeling and geometric Brownian motion modeling [31]. A related technique based on the analysis of short-term vs. long-term attainability and controllability has been followed by [32] and [33].

Retrospective signal analysis of emission changes becomes important when countries seek to assess their actual achievements in the commitment year/period. We distinguish between two fundamentally different approaches: static and dynamic. The static approach is identical to the one taken under preparatory signal analysis except that the agreed emission limitation or reduction commitment is replaced by the actual emission achievement. The emission signal is evaluated in terms of uncertainty, detectability or statistical significance, risk, etc. In contrast, the dynamic approach additionally considers how the emission signal has actually evolved between the base year and the commitment year/period, taking its dynamics into account. Here, expertise gained under midway analysis can serve as a platform as it also aims at evaluating full emission paths.

In their commensurability exercise, with its focus on six preparatory signal analysis techniques, Jonas and colleagues [23] concluded that a single best technique does not, and most likely will not, exist.⁴ This is because the available techniques

² It is noted that attempts exist to put one of these six techniques to analyze uncertain emission changes, the verification time concept, on a stochastic basis (see Ermolieva et al., herein; and also [27] and [28]). It is correct to say that this technique still undergoes scientific scrutiny and awaits adjustment in order to operate in a preparatory mode.

³ See http://www.iiasa.ac.at/Research/FOR/unc_overview.html for an overview on IIASA's monitoring reports and the countries that are monitored.

⁴ For the authors' study and numerical results see http://www.iiasa.ac.at/Research/FOR/unc_prep.html. Referring readers to this website facilitates easy replication for follow-up studies or, as in this case, avoiding duplication.

suffer from inconsistencies in dealing with uncertainty that are not scientific but that are related to the way the Kyoto Protocol was designed. One technique, e.g., allows a country with a smaller emission reduction commitment to gain an advantage over a country with a greater emission reduction commitment;⁵ while another technique forces countries a priori to realize detectable signals before they are permitted to make economic use of their excess emission reductions.⁶ Jonas and colleagues [23] stress that these “inconsistencies” are the consequence of the Kyoto policy process running ahead of science, leaving us with the awkward problem of choosing between bad or undesirable alternatives in applying preparatory signal analysis. We can simply ignore uncertainty knowing that, e.g., emission markets will then lack scientific credibility or we can give preference to one preparatory technique over another knowing that none is ideal in satisfying the Protocol’s political cornerstones. As two of the most important shortfalls on the side of policymaking the authors [23] identify (1) the overall neglect of uncertainty confronting experts with the finding that for most countries the agreed emission changes are of the same order of magnitude as the uncertainty that underlies their combined CO₂ equivalent emissions (compare the right column of Table 11.2 with the second column of Table 11.3); and (2) the existence of nonuniform emission limitation or reduction commitments that were determined “off the cuff” (i.e., derived via horse-trading) and did not result from rigorous scientific considerations.⁷ From a purely quantitative point of view, the first shortfall is of greater relevance than the second one.

⁵ See, for instance, the so-called undershooting (Und) concept: Excel file available via *numerical results* to [23] at http://www.iiasa.ac.at/Research/FOR/unc_prep.html: Worksheet *Undershooting 4*:

column C = Kyoto commitments δ_{KP} for country groups 1–8 (see also Table 11.2);
 column E = the accepted risk α that a country’s true emissions in the commitment year/period are equal to, or greater than, the country’s true Kyoto target (risk α can be grasped although true emissions and targets derived from them are unknown by nature); and
 columns F–N or U–AC (restricted to rows 14–16) = presumed relative uncertainty ρ of the country’s reported emissions.

The Und concept requires undershooting of the countries’ Kyoto targets in the commitment year in order to handle and decrease risk α (see columns F–N, rows ≥ 17 , for the required undershooting). Varying δ_{KP} while keeping the relative uncertainty ρ and the risk α constant (e.g., at $\rho = 15\%$ and $\alpha = 0.3$) exhibits that countries complying with a smaller δ_{KP} are better off than countries that must comply with a greater δ_{KP} (see columns U–AC, rows ≥ 17 , for the modified emission limitation or reduction target, which is the sum of the agreed target under the Kyoto Protocol plus the required undershooting). Such a situation is not in line with the spirit of the Kyoto Protocol!

⁶ See, for instance, the so-called combined undershoot and verification time (Und&VT) concept: Excel file available via *numerical results* to [23] at http://www.iiasa.ac.at/Research/FOR/unc_prep.html: Worksheet *Und&VT 1*: Fig. 1 therein. The Und&VT concept requires a priori detectable emission reductions, not limitations. That is, it requires the Protocol’s emission limitation or reduction targets to be corrected for nondetectability through the introduction of an initial or obligatory undershooting so that the countries’ emission signals become detectable before the countries are permitted to make economic use of their excess emission reductions. This nullifies, de facto, the politically agreed targets under the Kyoto Protocol!

⁷ The situation would be different if the nonuniformity of the emission limitation or reduction commitments would be the outcome of a rigorously based process resulting in a straightforward

However, it appears that the first shortfall will vanish soon as mankind is increasingly under pressure to adopt and realize greater emission reductions in the mid to long-term [36], [37, Decision 1/CP.13]. Notwithstanding, we would still be left with the problem of which analysis technique to give preference to. This discussion has not even started.

11.4 How to Deal with Uncertainty?

In this section we elaborate on Sect. 11.2 from a holistic point of view. Our starting point is FCA (or more generally, full accounting of all emissions and removals of all GHGs). We consider FCA a prerequisite for constraining and reducing uncertainties in our understanding of the global climate system. A dual-constrained (verified) full carbon analysis can compare the sum of Earth-based measurements of flows to and from the atmosphere with atmosphere-based evaluation of exchanges with the Earth. As specified by [5], a verified FCA, including all sources and sinks of both the technosphere and terrestrial biosphere considered continuously over time, would allow the research and inventory communities to:

- Present a real picture of emissions and removals at continental and smaller scales.
- Avoid ambiguities generated by such terms as “managed biosphere,” “base-line activities,” and “additionality.” Elimination of splitting the terrestrial biosphere into directly human-impacted (managed) and not directly human-impacted (natural) parts would be advantageous from a verification point of view as there is no atmospheric measurement that can discriminate between the two [21, Sect. 3].
- Make available reliable and comprehensive estimates of uncertainties that cannot be achieved using the current approach under the UNFCCC and Kyoto Protocol [8, 9]. It is impossible to estimate the reliability of any system output if only part of the system is considered. The tacit assumptions underlying the Protocol are that man’s impact on nature, the not-accounted remainder under the Protocol, is irrelevant and inventory uncertainty only matters from a relative point of view over space and time, not an absolute one. However, this approach is highly problematic because biases (i.e., discrepancies between true and reported emissions), typically resulting from partial accounting, are not uniform across space and time. In addition, man’s impact on nature need not be constant or negligible.⁸

rule that applies equally to all countries as would be the case, for instance, under the so-called contraction and convergence approach (e.g., [34, Sect. 2.3.2], [35]).

⁸ In their recent study [38, Table 1] show that, making use of global carbon budget data between 1959 and 2006, the efficiency of natural carbon sinks to remove atmospheric CO₂ has declined by about 2.5% per decade. Although this decline may look modest, it represents a mean net “source” to the atmosphere of 0.13 PgC y⁻¹ during 2000–2006. In comparison, a 5% reduction in the mean global fossil emissions during the same time period yields a net “sink” of 0.38 PgC y⁻¹. Thus, deteriorating natural carbon sinks as a result of climate change or man’s direct impact exhibit the potential to offset efforts to reduce fossil fuel emissions. This shows that man’s impact on nature

FCA is essential for good science. However, it would be for policymakers to decide how FCA is used. FCA could be used for “crediting” in the sense of the Kyoto Protocol (i.e., for compliance) or only for “accounting” and scientific understanding as required under the UNFCCC. Given that the treatment of the land use, land-use change, and forestry (LULUCF) sector in general poses a number of characteristic challenges (see box), we prefer FCA accounting under which, however, the Kyoto compliance accounting as required under the Protocol would form a logical and consistent subset.

Uncertainty in the LULUCF sector

Expressing uncertainties in the LULUCF sector can be challenging because of:

- The complexities and scales of the systems being modeled.
- The fact that human activity in a given year can impact emissions and removals in these systems over several years, not just the year in which the activity took place.
- These systems being strongly affected by inter-annual, decadal, and long-term variability in climate.

Knowledge of the temporal dynamics of systems – what has happened in the past, and how actions in the present will affect emissions/removals in the future – is important; gaps in this knowledge add to uncertainties about the immediate impacts of human activities.

Approaches to estimating emissions and removals in the LULUCF sector frequently involve the use of detailed data and computer models to simulate the complex functional relationships that exist in natural systems. But a consequence of using more detailed methods is that the estimation of uncertainty also comes more into play. However, despite conceptual and technical challenges, powerful tools for combining different kinds of information from multiple sources are becoming available and are increasingly being used by modelers to reduce uncertainties in the LULUCF sector.

These tools allow modelers to increase their focus on model validation and on reconciling results from alternative approaches. However, one key barrier remains. Reporting under the UNFCCC and Kyoto Protocol provides only a partial account of what is happening in the LULUCF sector. To close the verification loop would require the adoption of FCA.

Despite improvements in approaches to estimating uncertainty in emissions and removals in the LULUCF sector, some challenges remain. The treatment of this sector in future policy regimes requires special consideration.

is indeed not negligible and stresses the need to look at the entire system, that is, to develop a FCA system where emissions and removals and their trends are monitored in toto.

FCA is expected to facilitate the reconciliation of two broad accounting approaches: top-down and bottom-up accounting. Top-down accounting takes the point of view of the atmosphere. It relies on observations of atmospheric CO₂ concentrations, changes in concentrations, and atmospheric modeling to infer fluxes from land and ocean sources. Bottom-up accounting takes the opposite perspective. It relies on observations of stock changes or fluxes at the Earth's surface and infers the change in the atmosphere. FCA – estimating all land and ocean-based fluxes, whether human-induced or not – is necessary to reconcile the top-down and bottom-up approaches.

While methods of both top-down and bottom-up accounting have improved in recent years, both approaches still have areas of weakness. Investment in research is needed to tackle these limitations, improve the FCA approach, and hence reduce uncertainties (see also [39]).

Last, but not least, it must be kept in mind that verification of emission estimates does not necessarily imply detection of emission signals (e.g., decreased emissions) over time. It is the detection of emission signals that is needed to complement the bottom-up/top-down accounting of GHGs and the prime goal of this research community to close the existing gaps in the accounting.

Thus, from a policy perspective, there are pressing issues regarding how uncertainty can be dealt with through uncertainty analysis techniques and improvements to science. The implications for policymakers working to reduce human impacts on the global climate include [5]:

- Uncertainty analysis helps to understand uncertainties: better science helps to reduce uncertainties. Better science needs support, encouragement, and investment. FCA in national GHG inventories is important for advancing the science. It could be included in reporting but separated from targets for reducing emissions.
- Uncertainty is inherently higher for some aspects of an inventory than for others. For example, the LULUCF sector has higher uncertainties than the fossil fuel sector and estimates of N₂O emissions tend to be more uncertain than those of CH₄ and CO₂. This raises the possibility that in designing future policy agreements some components of a GHG inventory could be treated differently than others.⁹
- Improving inventories requires one approach; improving emissions trading mechanisms another. Inventories will be improved by increasing their scope to include FCA. In contrast, one option for improving emissions trading mechanisms would be to reduce their scope. Currently, emissions trading mechanisms may include estimation methodologies with varying degrees of uncertainty but they do not explicitly consider uncertainty or treat it in a standardized fashion. There are two options for improving this situation. The first option, as mentioned, is to reduce

⁹ This view of treating subsystems individually and differently runs counter to the approach typically taken. The tendency has been to treat subsystems collectively and equally and to dispose over a wide range of options in order to minimize costs or maximize benefits resulting from the joint emissions reduction of GHGs and air pollutants (e.g., [40], [41, (77)], [42]).

the scope of emissions trading mechanisms – by excluding uncertain methodologies or more uncertain GHGs – to make them more manageable (see also [43]). The second option is to retain the scope of emissions trading mechanisms but to adopt a standardized approach to estimating uncertainty. But we could not guarantee that the latter approach would eventually withstand large biases resulting, e.g., from a mismatch in the bottom-up vs. top-down accounting.

In the context of pricing uncertainty, it needs to be mentioned that uncertainty is an inherent part of any emissions accounting and that it will play an important role in both the scientific understanding of emissions and in their political treatment. At present, however, uncertainty does not play a role in trading of emissions credits. Ultimately, uncertainty can be borne by either the buyer or seller of any asset, and it should be agreed in advance of any exchange how this is to be dealt with. Risky or uncertain assets will be traded at a discount to the extent that the risk and uncertainty are to be assumed by the buyer.

Literature on treating scientific uncertainty upfront in financial markets is already emerging (e.g., [26, 44, 45]), but this has not yet been applied widely to GHG emissions credits. For now it appears that buyers of emissions credits generally accept credits without uncertainty and the seller is obligated to ensure that the credits are fulfilled.

With the current system of trading in allowances and credits, neither buyers nor sellers have much incentive to reduce the uncertainty associated with emissions inventory or reduction estimates; to do so might impact the single-point emission (or reduction) estimate, thus directly affecting the value of allowances or credits. For example, a highly uncertain emission reduction estimate that is biased high will tend to be worth more (claiming greater reductions), presupposing the market's willingness-to-pay, than the same reduction figured more accurately and with greater uncertainty. This suggests the possibility that other, more complex, pricing mechanisms than the current cap-and-trade system might exist and would be better able to deal with uncertainty by, for example, monetizing (i.e., rewarding) increased confidence. Such a system might also differentiate between different types of emissions and/or reductions and their uncertainties.

11.5 Conclusions

We see a clear rationale for conducting and improving uncertainty analysis:

- Uncertainty analysis improves the monitoring of GHG emissions. Uncertainty analysis helps to understand uncertainties and encourages better science that will help to reduce uncertainty.
- Better science requires the adoption of FCA. More investment in research is needed to reconcile the bottom-up and top-down accounting approaches that are fundamental to FCA and that dual-constrain uncertainty. FCA may only be used for “accounting” but with the Kyoto compliance accounting as a logical

and consistent subset used for “crediting.” We anticipate that within a few years scientists will overcome still existing bottom-up/top-down accounting gaps for the Kyoto GHGs at the scale of continents. Scientists may even be able to down-scale validated, and verified (dual-constrained), emissions estimates to the scale of countries or groups of countries. That is, scientists will be able to verify (correct) politically driven (mis-)accounting reported annually bottom-up under the Kyoto Protocol and its successor.

- Some GHG emissions and removals estimates are more uncertain than others. Options exist to address this issue, and these could be incorporated in the design of future policy regimes. These options also include (1) the option of not splitting the terrestrial biosphere into a directly human-impacted (managed) and a not-directly human-impacted (natural) part to avoid sacrificing bottom-up/top-down verification; and (2) the option of not pooling subsystems, including sources and sinks, with different relative uncertainties but treating them individually and differently.
- We expect the treatment of scientific uncertainty in emission trading markets to gain relevance. Neither buyers nor sellers of GHG emission credits have a strong interest to let this issue go unresolved.
- The issue of compliance also goes unresolved and requires directing attention to the appropriate treatment of emission changes in consideration of uncertainty. Currently, signal analysis is still treated independently of bottom-up/top-down verification, but scientists will eventually be able to make the two consistent and to go hand-in-hand.

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Chapter 12

Uncertainty Analysis of Weather Controlled Systems

K.J. Keesman and T. Doeswijk

Abstract The indoor climate of many storage facilities for agricultural produce is controlled by mixing ambient air with the air flow through the store room. Hence, the indoor climate is affected by the ambient weather conditions. Given hourly fluctuating energy tariffs, weather forecasts over some days are required to effectively anticipate. Hence, typically a real-time optimal control strategy results. As weather forecasts are uncertain, predicted model outputs and related costs of the control strategy become uncertain. Usually, a medium-range weather forecast for a period of some days consists of an ensemble of forecasts. Hence, the uncertainty in the weather forecast is known a priori. In addition to this, in past-performance studies where weather forecasts and observed weather variables are given, an a posteriori evaluation of the forecast errors can be made as well. The objective of this study is to evaluate the uncertainty in the costs related to weather forecast errors and uncertainty, given the control inputs. In a simulation case-study with real weather forecasts and observed weather, it appeared that only slight cost increases can be expected due to errors and uncertainties in weather forecasts if the optimal control problem is calculated every 6–12 h in a receding horizon context.

12.1 Introduction

Indoor climate in greenhouses, office buildings and storage facilities for agricultural produce are generally affected by outdoor weather conditions [2, 9, 13], for instance by heat transfer through the boundaries, solar radiation and ventilation with outdoor air. In addition to this, the indoor climate is also affected by the respiration and evapotranspiration of living creatures and biological material, or by active heating or cooling. Typically, in practice, indoor temperatures and relative humidities are

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controlled by feedback controllers with P or PI structure (see [14] for an overview) using direct measurements of temperature and relative humidity. In this study, we focus on indoor climate control that uses ambient air in the air flow through the (store) room. Hence, optimal control strategies that are able to anticipate on future changes of the weather conditions or on hourly-daily changes in energy tariffs allow, in principle, a better cost-effective performance. Clearly, in such control strategies weather forecasts become important and uncertainties in weather forecasts then lead to uncertain predictions of the indoor climate. Moreover, the cost function related to any kind of optimal control becomes subject to uncertainties and errors in the weather forecasts. The sensitivity of the model outputs and the related costs therefore needs to be investigated. In what follows, we restrict ourselves to a so-called receding horizon optimal control (RHOC) strategy in discrete-time. The basic idea behind RHOC is that a finite-time optimization problem over the horizon H , with or without inequality constraints, is solved, i.e.

$$u(k, k + H) = \arg \min \phi(x(k + H)) + \sum_{\kappa=k}^{k+H} \mathcal{L}(x(\kappa), u(\kappa), d(\kappa)) \quad (12.1)$$

$$\text{s.t. } x(k + 1) = f(x(k), u(k), d(k)), \quad g(x(k), u(k)) \leq 0$$

$$x_0 = x(k)$$

Subsequently, only the first $l < H$ inputs are implemented and at $k + l$, given an update of x_0 , the optimization problem is solved again, but now on the time interval $[k + l, k + l + H]$. For both air conditioned buildings [7] and potato storage facilities [8, 10] it was shown that a good weather forecast reduces the cost function almost as much as a perfect weather forecast. In these studies short term weather forecasts (1–2 days ahead) were used. However, because of the slow dynamics of products in storage facilities, knowledge of medium-range predictions of the indoor climate may be profitable. Hence, this study focuses on the effect of errors and uncertainties in medium-range (up to 10 days ahead) weather forecasts on the indoor climate predictions, in real-time, and associated product temperature and costs.

Nowadays, the medium-range weather forecast [11] consists of an ensemble of 50 different weather forecasts. All 50 ensemble members have an equal probability of occurring. Hence, the uncertainty, or variation, in the weather forecasts is known a priori. This knowledge can then be used to evaluate a calculated optimal control solution by calculating the differences in costs related to each of the ensemble members. Next to this, with observed weather given, in a post-performance analysis the optimal control solution can also be evaluated a posteriori.

The objective of the study is to evaluate the effect of errors and uncertainties in the weather forecasts on the costs of the calculated optimal control problem of a (potato) bulk storage facility.

This chapter is structured as follows: first a brief description of the bulk storage model and the cost function is given. Next, some information about the weather forecasts is provided. Then, open loop and closed loop evaluations using weather forecasts and observations are given in subsequent sections. Finally, the results are discussed and conclusions are drawn.

12.2 Preliminaries

12.2.1 Bulk Storage Model

Models that describes the dynamics of the product in a bulk storage facility can be found in, e.g. [6, 8, 9, 15–19]. The model developed in [9] was simplified to make it suitable for use in an RHOC algorithm [10]. A brief description of this discrete-time dynamic model is presented in this subsection. The model is given by

$$x_T(k+1) = x_T(k) + p_1 + [d_1(k) - x_T(k)] \left(p_3 + p_2 p_5 u_1(k) [p_7 + (1 - p_7) u_2(k)] \right) \quad (12.2)$$

$$x_{\text{CO}_2}(k+1) = x_{\text{CO}_2}(k) + p_1 p_4 + [d_2(k) - x_{\text{CO}_2}(k)] \left(p_2 u_1(k) [p_7 + (1 - p_7) u_2(k)] + p_6 \right) \quad (12.3)$$

where x_T represents the product temperature, x_{CO_2} the CO_2 concentration in the bulk, $d_1 = T_{wb,ext}$ the ambient wet bulb temperature, d_2 the ambient CO_2 concentration, $u_1 = u_{mix}$ the fraction of ambient air in the air flow, $u_2 = u_{vent}$ the fraction of maximum possible internal ventilation, and p a vector containing physical and design parameters. Hence, the control input is defined by: $u = [u_1, u_2]^T$. The sample time used for this model is 1 h.

The temperature of the bulk can be measured and thus (12.2) is regularly updated. The CO_2 -concentration, however, is difficult to measure in practice. The store room is, therefore, basically controlled on the product temperature. In the model-based RHOC strategy, the CO_2 -concentration is calculated using (12.3) without any measurement correction. The CO_2 -concentration is taken into account via constraints to avoid too high concentrations that lead to damage of the product. This approach suffices, because of the fast dynamics of the CO_2 -concentration, when the room is ventilated. That is, during ventilation the CO_2 -concentration quickly reaches the known external CO_2 -concentration, so that large modeling errors are avoided. Consequently, right after a period of ventilation, $x_{\text{CO}_2}(k)$ is equal to $d_2(k)$.

12.2.2 Weather Forecasts

The medium-range weather forecasts have been provided by Weathernews Benelux. These forecasts contain hourly forecasts of weather variables up to 10 days ahead. The ensemble weather forecasts used in this paper consist of 50 ensemble members with equal chance of occurring. In what follows, the ensemble mean is used as the nominal weather forecast.

New weather forecasts and actual weather observations become available every 24 and 6 h, respectively. It has been shown in [3] that short-term weather forecasts

(up to 36 h ahead) can be improved by using more frequent, local observations in combination with Kalman filtering techniques. A similar method is used here to correct the medium-range weather forecasts every 6 h.

12.2.3 Cost Function

To give some physical insight into the control objectives of the potato storage facility, the elements of the cost function are mentioned below. How the weighting factors in the cost function are chosen is beyond the scope of this paper. In our application, the following control objectives are to be fulfilled:

- The temperature of the bulk must be kept as close as possible to a pre-specified reference temperature T_{ref} (i.e. minimize $\|x_T(k) - T_{ref}\|$).
- The temperature must always be kept above a specified minimum temperature T_{min} (i.e. inequality constraint: $x_T(k) > T_{min}$)
- The temperature may not decrease faster than a specified limit T_{Δ} within 24 h (i.e. inequality constraint: $x_T(k - 24) - x_T(k) < T_{\Delta}$).
- The weight loss of the product due to evaporation must be as small as possible (i.e. minimize $\sum_{k=0}^H f_1(x_T(k), u(k), d(k))$ with H the prediction horizon.
- The CO_2 -concentration must always be kept below a specified maximum $\text{CO}_{2,max}$ (i.e. inequality constraint: $x_{\text{CO}_2} < \text{CO}_{2,max}$)
- The energy costs related to ventilation must be as small as possible (i.e. minimize $\sum_{k=0}^H f_2(u_2(k))$).

12.2.4 Receding Horizon Optimal Control

Given the model (12.2)–(12.3) with corresponding initial states, a weather forecast containing $d_1(k)$ for $k = 1, \dots, H$ and assuming $d_2(k) = 0.0314$ to be constant, control trajectories of u_1 (fraction of ambient air in the air flow) and u_2 (fraction of maximum possible internal ventilation) can be calculated such that a cost function on the time interval $[0, H]$ is minimized according to:

$$\min_u J = \min_u \phi(x(H)) + \sum_{k=0}^H \mathcal{L}(x(k), u(k), E[d(k)]) \quad (12.4)$$

where the expected value (denoted by $E[\cdot]$) is taken because the weather forecast is a stochastic variable. In our application of bulk storage in a store room there are no final costs at the end of the prediction horizon, i.e. $\phi(x(H)) = 0$. From the preceding subsection we notice that $\mathcal{L}(\cdot)$ contains a weighted combination of $\|x_T(k) - T_{ref}\|$, $f_1(\cdot)$ and $f_2(\cdot)$. If this open-loop control problem is solved repeatedly every l hours (with $l < H$) given the updated (or measured) states

and weather forecasts the control loop is closed. As mentioned before, this type of control strategy is called receding horizon optimal control (RHOC).

In the following, the RHOC solution with the nominal weather forecast is taken as the reference point for the uncertainty evaluation. For the evaluation a period from 13 April to 2 June 2005 (almost 51 days) had be chosen. The medium-range weather forecasts have been obtained for location “De Bilt, The Netherlands”. The forecast horizon of the weather forecast also determines the maximum horizon of the RHOC strategy ($H = 217$ h). In fact, one would expect a maximum horizon of $(10 \times 24 + 1)$ h. However, the medium-range forecasts are released with a delay of 1 day and thus $H = 241 - 24 = 217$ h. During the evaluation period every 6 h a new optimal control trajectory was calculated (i.e. $l = 6$) with an updated weather forecast, according to the procedure suggested in [3]. For each optimal control run the initial conditions of the states were set to the corresponding measured values. In an RHOC framework the calculated optimal inputs for the pre-specified control interval (l) of 6 h are implemented and recalculated when after 6 h new information becomes available. Consequently, in our application the optimal control input trajectories are calculated over an interval of 217 h and updated every 6 h. In total 203 optimal control trajectories, given the nominal weather forecasts, are calculated (four times a day, almost 51 days). An example of a calculated control trajectory with the accompanying predicted state and cost evolution is presented in Fig. 12.1. The reference temperature (T_{ref}) in this case was 7°C . Notice that the calculated open loop optimal control inputs are in antiphase with the external wet bulb temperature. As expected, the product temperature (T_p) follows the ventilation pattern. Notice, furthermore, that with increasing temperature differences between the product temperature and the external wet bulb temperature also the costs increase accordingly.

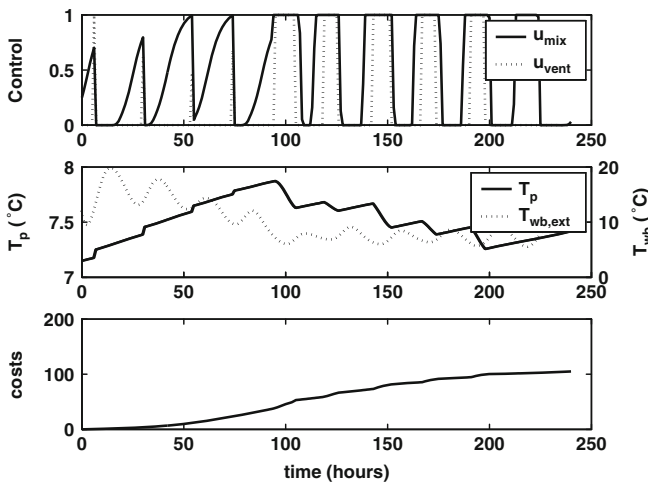


Fig. 12.1 Optimal control output starting from 1 May 2005. The sub-figures show respectively: calculated optimal controls; predicted product temperature and external wet bulb temperature; predicted cumulative costs

12.3 Weather Forecast Uncertainty and Error Analysis

In this section, we investigate several possibilities to evaluate the weather forecasts errors and uncertainties on the costs. First, the change in costs is investigated when observations of the weather are used instead of the nominal forecast. Hence, this analysis, based on 203 control trajectories with associated costs, evaluates the effect of errors in the weather forecasts on the costs. Second, the effect of individual ensemble members on the costs is investigated. Thus, this type of analysis evaluates the maximum effect of variation in the ensemble on the absolute costs.

12.3.1 Open Loop Evaluation

Let us start by evaluating the effect on the absolute costs when observed instead of nominal forecasted weather data is used. Recall that the prediction horizon is 217h, which defines the maximum length of the evaluation. However, in what follows the summation variable r is introduced to account for intermediate changes in the costs. If now the model with fixed optimal control trajectories is ran again but with observed weather (d_1 and d_2) instead of nominal forecasted weather data, a change in costs is observed. In Fig. 12.2, 203 differences between calculated (using forecasts) and realized (using observations) running costs, as a function of the summation variable r , are presented. Herein, the running costs are defined by:

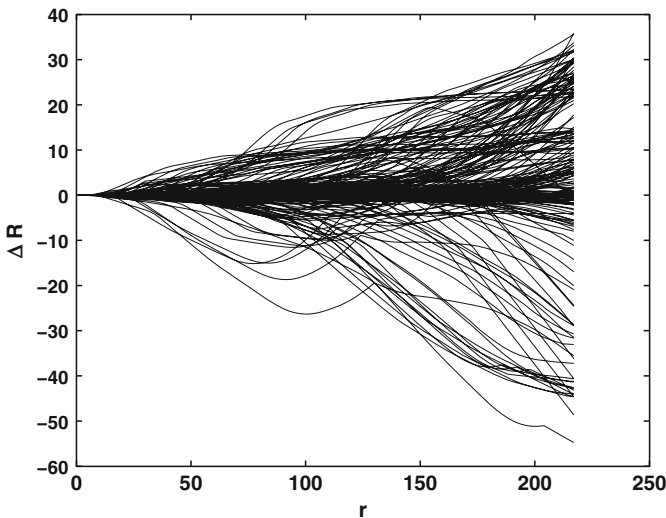


Fig. 12.2 Difference in calculated absolute costs and realized costs for each of the 203 optimal control runs

$$R(r) = \sum_{k=0}^r \mathcal{L}(x(k), u(k), d(k)) \quad \text{for } r = 1, \dots, H \quad (12.5)$$

For what follows, we define $J \triangleq R(H)$.

From Fig. 12.2, it can be seen that the total cost difference, as expected, can be both positive and negative. This implies that, given the optimal control trajectory based on forecasts, the realized weather can reduce the total costs more than was expected from the forecasted weather. However, it does not mean that, if the observed weather was used in the RHOC calculations (which is only possible afterwards), the calculated optimal control trajectory generates minimal costs. Furthermore, histograms derived from Fig. 12.2 for different r show that the mode is around $\Delta R = 0$, and that the frequency distributions are rather skewed with a “thin” tail for ΔR negative.

Since the total absolute costs J change for every optimal control run, because of changing initial states and changing weather forecasts after each 6 h, the relative change in the costs, i.e. $\Delta R^{rel} = \frac{R_{obs} - R_{fct}}{J_{obs}}$, is calculated as well and presented in Fig. 12.3. In addition to the absolute or relative differences, we noticed that the absolute costs R can change dramatically (not shown here) over the evaluation period from 13 April to 2 June, because the outdoor temperature significantly increases. Especially, the cost criterion term $\|x_{k,T} - T_{ref}\|$ then increases. Relatively to the costs at the beginning or the end of the prediction horizon H , however, the change in costs does not seem to change that dramatically.

Using the observed weather to calculate the costs is a useful a posteriori tool, since it evaluates the positive and negative costs when controlling a system under expected disturbances. Nevertheless, it should be emphasized that the conclusions

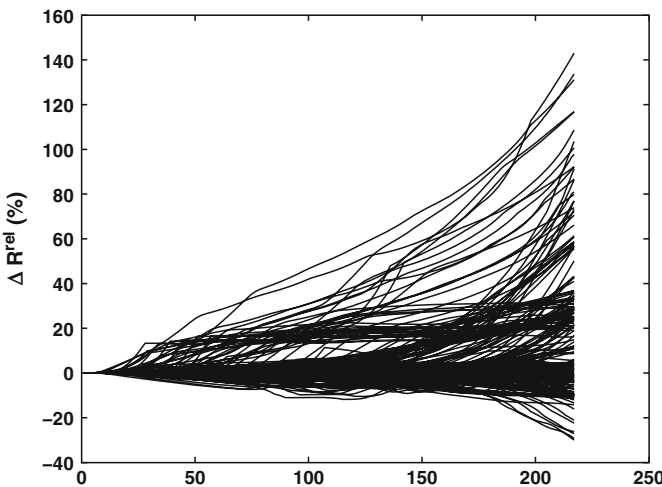


Fig. 12.3 Difference in calculated relative costs and realized costs, for $r = 1, \dots, H$, for each of the 203 optimal control runs

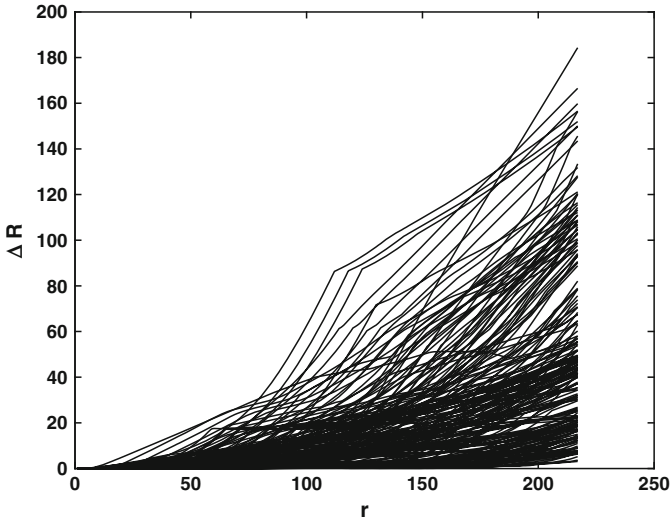


Fig. 12.4 Difference in calculated absolute costs and maximum costs based on the ensemble related to each of the 203 optimal control runs

certainly depend on the weather data and thus the type of weather. For a full analysis of the posterior behavior, other periods over a couple of years should have been evaluated, as well. However, such analysis was out of the scope of this study. To evaluate the uncertainty of the costs a priori, however, other information about the uncertainty of the weather forecast is needed. As mentioned before, the uncertainty of the weather forecast is embedded within the ensemble (see Sect. 12.2.2). By calculating the costs related to each of the ensemble members the worst case scenario can be evaluated, i.e. the ensemble member for which the costs are highest of all for $r = H$. In Fig. 12.4 the worst-case cost differences are presented, again as a function of r with $r = 1, \dots, H$. Since only the worst case differences, related to the largest difference at $r = H = 217$, are presented, only 203 out of 50×203 runs are shown. It can be seen here that the worst-case scenarios always lead to increased costs, as expected, and these costs are considerably larger than the realized costs (Fig. 12.2). The question remains what the effect of a receding horizon control strategy on the uncertainty in costs is. This question will be answered in the next subsection.

12.3.2 Closed Loop Evaluation

In the specific RHOC implementation every 6 h, when actual weather data and updated forecasts become available, the optimal control trajectories are recalculated from (12.4) and implemented. Hence, only the control inputs for the first 6 h are

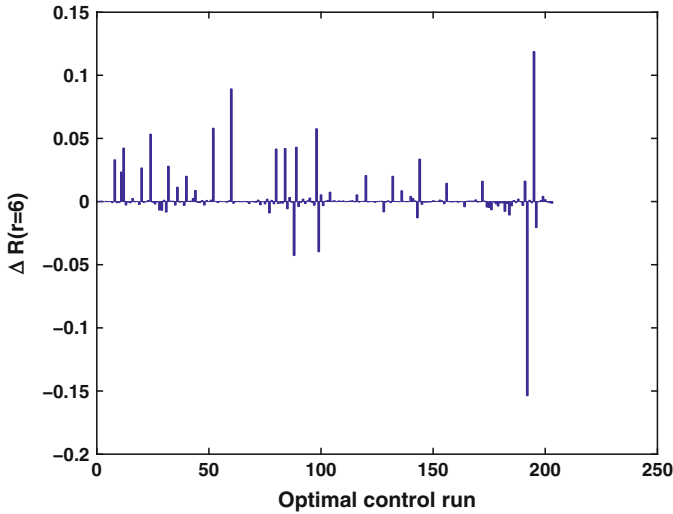


Fig. 12.5 Difference in calculated costs and realized costs after 6 h for each of the 203 optimal control runs

effectively used in the control strategy. This also implies that the uncertainties and errors in the cost function after 6 h needs to be evaluated. In Fig. 12.5 the differences between calculated costs, after 6 h because of the updates and using nominal weather, and actual costs based on observed weather are presented for each of the 203 control runs. Hence, this allows an a posteriori evaluation of the forecast errors. The small differences in costs after 6 h can be clearly seen from this figure. Notice that the small magnitude of the differences could also have been seen from Fig. 12.4. If all the costs shown in Fig. 12.5 are summed the additional costs due to errors in the nominal weather forecasts are known for the given period from 13 April to 2 June 2005.

From the a priori known weather forecast uncertainties, which can be directly obtained from the ensemble predictions, the worst-case scenario can be calculated as before. It appeared that the differences in calculated costs and maximum costs after 6 h (not shown here) based on the ensemble related to each of the 203 optimal control runs are very small, $\Delta R(r = 6) < 0.2$. However, the question remains how ΔR will evolve when the control horizon ℓ is chosen larger. In Table 12.1 the maximum (relative) deviation in the cost function, obtained from the weather forecast ensemble, is given for different control intervals (l).

Hence, for this case study it can be concluded that applying feedback every 6 h will reduce the uncertainty in the calculated costs, due to weather forecast errors and uncertainty, tremendously. It can also be seen from Table 12.1 that the difference in costs up to 24 h remains relatively small. In other words, in case of a communication failure between the optimization algorithm and the control computer of less than, e.g. 1 day no manual intervenience is required. However, it should be realized that these statements are only valid for our specific application in the pre-specified

Table 12.1 Additional costs of the realized costs and the worst-case scenarios for weather forecast uncertainty

ℓ	$\Sigma(\Delta J)/J$ (%)
6	0.37
12	1.15
18	2.05
24	3.17
48	8.0
72	14.2

evaluation period. Nevertheless, as can be seen from Figs. 12.2–12.4, the weather forecasts do not show much variation in the first 12 h. This phenomenon has been observed for other periods, as well. Thus, for $l \leq 12$, we conclude that in general the effect of the weather forecast uncertainty on the costs, as defined in (12.4), is marginal. Hence, under the assumption that CO₂ is never limiting, a control horizon of $l = 12$ instead of 6 would suffice in practice. Clearly, the weather forecast ensemble can be used for an a priori uncertainty evaluation on the costs. But we would be more flexible if we could anticipate on the forecast uncertainty. This idea will be further exploited in the next section.

12.4 Discussion

If in open loop the calculated costs based on the weather forecast and the *actual* costs, given observed weather and realized controls, are compared, it is evident that the costs can both increase and decrease. Not only the total costs are of importance but also the relative costs. During a long warm period the calculated costs will increase significantly. The absolute increase or decrease due to uncertain weather forecasts can then be quite large whereas the relative change may be reasonable. The opposite may also occur, i.e. large relative change versus a low absolute change.

Prior knowledge about the uncertainty of the forthcoming weather is very useful to study uncertainties in the near-future costs. From Figs. 12.2 and 12.4 it can be seen that, for this specific case, the (expected) cost increase is in general much larger for the a priori forecast uncertainty than for the a posteriori forecast error. Similar conclusions can be drawn in the closed loop case, see Fig. 12.5. From these figures it can be seen that the ensemble, indeed, is a useful tool to evaluate the uncertainty of the cost function a priori.

Recall that in optimal control theory a cost function, like (12.4), is minimized by adjusting the control inputs u , see, e.g. [12]. In case of the storage facility, genuine optimal control trajectories could be calculated if the (near) future weather would be exactly known in advance. However, future weather forecasts are never exactly known. A common approach, as we introduced in the previous subsections, is to define a cost function on the basis of the nominal disturbance trajectory (d^0 , i.e. the nominal weather forecast), solve the optimization problem and evaluate the uncertainties in the costs. Alternatively, an approach based on implicit formulation of uncertainties in the cost function could have been chosen, such that an

effective uncertainty reduction can be obtained. A classical example of this approach is minimum variance control [1].

For instance, in the case of our storage facility, minimum variance control would lead to $u = 0$, that is no ventilation of the store room, because the uncertainty in the product temperature is directly related to the weather forecast. Clearly, the product temperature uncertainty increases during ventilation. Hence, minimum variance control is not an option in our case. Thus, future work should focus on appropriate extensions of the cost function (12.4), for instance in line with [5] or [4].

12.5 Concluding Remarks

In post-harvest storage of agricultural produce, optimal control strategies can be used to anticipate on future weather conditions. In the simulation case-study with real weather forecasts and observed weather it appeared that there are only slight cost increases due to uncertainty and errors in weather forecasts if the optimal control problem is calculated every 6 h within a RHOC framework. In our application, even an increase of the control horizon to 24 h leads to a maximum increase of less than 5%.

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Chapter 13

Estimation of the Error in Carbon Dioxide Column Abundances Retrieved from GOSAT Data

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Abstract In this chapter, the estimation of error in the method used to retrieve carbon dioxide (CO₂) column abundances obtained from the Greenhouse Gases Observing Satellite (GOSAT) is presented. GOSAT will be the first satellite dedicated to primarily observe CO₂ and methane (CH₄), which are considered to be major greenhouse gases, from space. The column abundances of CO₂ and CH₄ can also be retrieved using IMG data and SCIAMACY data. The retrieved CO₂ and CH₄ column abundances can be evaluated using spectral fitting residuals. On the other hand, there are uncertainties in the retrieved column abundances due to the error in the input parameters, such as the temperature profile and the water vapor profile during spectral observation. The estimated error of the retrieved column abundances using IMG data and SCYAMACY data is not affected by the error in input parameters. In this study, the retrieval error due to input parameters using factorial experiments is described. Finally, some application results are given.

13.1 Introduction

Global environmental problems, such as global warming, desertification and ozone depletion, have recently become major concerns. The Intergovernmental Panel on Climate Change (IPCC) recently said that “it is *extremely unlikely* (<5%) that the global pattern of warming during the past half century can be explained without

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external forcing, and *very unlikely* that it is due to known natural external causes alone. Greenhouse gas forcing has *very likely* caused most of the observed global warming over the last 50 years” [13]. Water vapor is the most important greenhouse gas, followed by carbon dioxide (CO₂). Methane (CH₄), nitrous oxide (N₂O), ozone (O₃), and several other gases present in the atmosphere in small amounts also contribute to the greenhouse effect [13]. Gases such as CO₂, CH₄, N₂O, and O₃, which are present in low concentrations in the atmosphere (<0.03% [21]) are referred to as “trace gases”. Therefore, an understanding of the characteristics of CO₂ on a global scale is crucial to solving global environmental problems.

Observing trace gases that are distributed very sparsely across the Earth’s surface is difficult. Despite the fact that oceans account for roughly two-thirds of the Earth’s surface, there are very few observation stations located on the oceans. Thus, it is important to develop methods that allow the observation of trace gases over the Earth. Satellite remote sensing provides one of the most useful methods for global atmospheric observations despite several limitations. Firstly, satellites cannot record data if clouds obscure the sensor’s view. Secondly, observation time is determined based on the local time when a satellite’s orbit is sun-synchronous polar orbit. Thirdly, solar radiance and Earth irradiance at specific wavelengths change due to the effects of trace gases. Since the trace gas contribution to the spectrum is so small, it is extremely difficult to precisely detect the physical quantity of a trace gas using satellite remote sensing from the measured spectrum. Furthermore, the spectral difference occurs at very narrow wavelengths of the specific solar radiance bands. Therefore, more advanced techniques are required to obtain spectral data with high resolution and high sensitivity.

The Interferometric Monitor for Greenhouse gases (IMG) aboard the Advanced Earth Observing Satellite (ADEOS), launched in August 1996, was the first high resolution nadir infrared instrument allowing the simultaneous measurement from space of a series of tropospheric trace gases: H₂O, CO₂, N₂O, CH₄, CFCs, O₃, and CO₃ [1, 7]. IMG was designed to measure the terrestrial thermal radiation in an infrared spectrum at a high resolution. The Scanning Imaging Absorption Spectrometer for Atmosphere Cartography (SCIAMACY) onboard the European Environmental Satellite (ENVISAT) observes the atmosphere with moderate spectral resolution. An algorithm is currently being developed and refined primarily for the retrieval of CH₄, CO, CO₂, H₂O, N₂O, and O₂ columns using SCIAMACY data [5]. In Japan, the Greenhouse Gases Observing Satellite (GOSAT) will be launched in early 2009. In the US, the Orbiting Carbon Observatory (OCO) satellite will also be launched in late 2008. GOSAT and OCO will also perform passive nadir observations in the near-infrared spectral region, but they will be optimized for CO₂ observations. It is expected that GOSAT and OCO will improve knowledge of CO₂ surface fluxes. In the GOSAT project, a global map of CO₂ and CH₄ column abundances will be distributed to users. Given the scope of these aims, it is important to clarify the precision of the GOSAT’s CO₂ column data. I. Aben quantifies the effect of aerosols and thin cirrus clouds in the atmosphere, which is the largest error source for CO₂ measurements from space. This study presents a whole evaluation method. Two types of error for the retrieved column abundance data are considered. Errors

derived from the spectral fitting residuals (internal errors) are obtained analytically. For the evaluation of the retrieved column abundance derived from the IMG data and the SCIAMACY data, the error is primarily internal error. In this study, errors derived from input parameters (external errors), which are used in the retrieval stage, are analyzed by examining factorial experiments, which are mainly used for quality control.

The structure of this paper is as follows. In Sect. 13.2, trace gas observations from satellite sensors are introduced, and the role of GOSAT in trace gas measurement is presented. The GOSAT mission is then described, and the previous error analysis by satellite remote sensing is presented. In Sect. 13.3, an evaluation method of GOSAT's CO₂ column abundance data is described in detail. Certain measurement geometry and atmospheric conditions are set, and the retrieval precision results are presented as one example. The results presented in this paper are not officially used but are referred to as the basic concepts in the GOSAT project, since the retrieval algorithm of the operational data processing is currently in the coding and tuning stage. The practical retrieval precision shall be made public at a future date. Section 13.4 concludes this paper.

13.2 Trace Gas Measurement by Satellite Remote Sensing

Trace gas measurement using satellite remote sensing is presented in Sect. 13.2.1, and the role of GOSAT in trace gas measurement from space is explained in Sect. 13.2.2. Lastly, the current evaluation methods of error in retrieved trace gas amounts are presented in Sect. 13.2.3.

13.2.1 *Observations of Trace Gases with Various Sensors*

Passive sensors measure solar radiance directly from the sun or reflected from the Earth. Both methods detect physical quantities of trace gases based on the differences in the spectrum observed from the trace gases in the atmosphere. Solar occultation is a technique that measures direct sunlight using limb observation of the Earth from the satellite. Observing the spectrum reflected from the Earth's surface is called "down-looking observation". It is necessary to use sensors with a high signal-to-noise ratio (S/N) and high spectral resolution. Advances in sensor technology combined with diffraction grating have allowed a Fourier Transform Spectrometer (FTS) system to detect the physical quantities of trace gases. For example, solar occultation sensors include the Improved Limb Atmospheric Spectrometer (ILAS) onboard the ADEOS satellite and the Stratospheric Aerosol and Gas Experiment (SAGE) and Halogen Occultation Experiment (HALOE) onboard the Upper Atmosphere Research Satellite (UARS). Solar occultation observations detect trace gases in the stratosphere at or above a height of 10 km. Below an altitude of 10 km, trace

gas values are not observed precisely, and the horizontal resolution is quite low. Moreover, since concentrations of CO₂ in the stratosphere are lower than in the troposphere, it is difficult to ascertain global CO₂ distributions using solar occultation technology.

Down-looking observations of trace gases are retrieved by both FTS and grating sensors. The Infrared Interferometer Spectrometer (IRIS) onboard the NIMBUS-3 satellite launched at 1969 was the first FTS type sensor. The IRIS instrumentation was a modified version of a classical Michelson Interferometer, and measures radiance in narrow (5 cm⁻¹) spectral intervals from 400 to 2,000 cm⁻¹ (5–25 μm) [9]. Two portions of the spectrum are used: 11 intervals each within the CO₂ band at 667 cm⁻¹ and the atmospheric window at 900 cm⁻¹. However, when NIMBUS-3 was launched, IRIS spectral resolution was lower than the width of trace gas absorption. Thus, it was considered that nadir observation of the tropospheric constituents was difficult, and that observation of greenhouse gases was not a significant objective.

ENVISAT with SCIAMACHY was launched in March 2002 in a sun-synchronous polar orbit. SCIAMACHY is outfitted with a grating spectrometer for measuring solar radiation reflected off the surface of the Earth, backscattered from the atmosphere, transmitted through the atmosphere, or emitted from the atmosphere in the ultraviolet, visible, and NIR spectral regions (240–1,750 nm, 1,940–2,040 nm, 2,265–2,380 nm) at moderate spectral resolutions (0.2–1.4 nm) [6]. CH₄, CO, CO₂, N₂O, and O₂ are retrieved from the SCIAMACHY near-infrared and visible spectra [5]. The standard deviation of the dry air column averaged mixing ratio XCO₂ within 1° latitude bands is ±10 ppmv or ±2.7% for measurements over land. The spatial (horizontal) resolution of SCIAMACHY nadir measurements depends on the orbital position and the spectral interval. The ground pixel (footprint) size is approximately identical at each orbital position. For example, the footprint is 60 × 30 km² for an integration time of 0.25 s, and 120 × 30 km² for 0.5 s.

The IMG sensor is a Fourier Transform type infrared spectrometer (FTIR) that was developed to measure greenhouse gases. The ADEOS satellite was in a near-polar sun-synchronous orbit, a 41-day recurrent polar orbit. IMG was developed for measuring greenhouse gases in the atmosphere, particularly in the troposphere, and was designed to measure the terrestrial thermal radiation in the infrared spectral region of 640–3,000 cm⁻¹ at a high resolution of 0.1 cm⁻¹ after reaching apodosis [12]. The footprint size is 8 km × 8 km, and the main atmospheric constituents, H₂O, CO₂, O₃, N₂O, CO, CH₄, and HNO₃ were retrieved. Error bars for the retrieved CO₂ at 1,000, 500, and 200 hPa, which are recognized as the 1-standard deviation (1-sigma) for analysis, are 6.03%, 3.93%, and 3.09%, respectively [14].

13.2.2 GOSAT Mission

The GOSAT project is a joint project of the Japan Aerospace Exploration Agency (JAXA), the Ministry of Environment (MOE), and the National Institute for Environmental Studies (NIES) [8,22]. JAXA is responsible for the satellite, its instrument

Table 13.1 Overview of GOSAT and orbit

Parameter	Value
(a) Body	
Size	Main body: 3.7 m (height) \times 1.8 m (width) \times 2.0 m (depth)
@@	Include solar paddle: 13.7 m
Weight	1,750 kg
Electric power	3.5 kW
Life span	5 years
(b) Orbit	
Orbit	Sun-synchronous polar orbit
Altitude	666 km
Inclination	98.05°
Period of revolution	14.66 orbits/day
Equatorial crossing time	Nominally 1 p.m. (\pm 15 min) local time (descending node)
Repeat coverage	3 days

development, and its operation, while MOE is involved in instrument development, and NIES is responsible for satellite data retrieval and data distribution. GOSAT will be launched in early 2009. An overview of the GOSAT satellite and the orbit is presented in Table 13.1.

Higher retrieval precisions are expected when the gas amount information is retrieved from the optimal wavelength band for the gas. The GOSAT mission will make the first global, space-based measurements of CO₂ with the precision, resolution, and coverage needed to characterize CO₂ sources and sinks at the regional scale. GOSAT carries two sensors: TANSO-FTS (Thermal And Near infrared Sensor for carbon Observation – Fourier Transform Spectrometer) that obtains solar radiance reflected from the ground surface; and TANSO-CAI (Thermal And Near infrared Sensor for carbon Observation – Cloud and Aerosol Imager) that obtains cloud and aerosol information of observed areas of TANSO-FTS. TANSO-FTS, which detects the gas absorption spectra of the solar short wavelength infrared (SWIR) reflected on the Earth's surface, as well as of the thermal infrared (TIR) radiated from the ground and the atmosphere. TANSO-FTS-SWIR data is obtained for three spectral bands, 0.76, 1.6, and 2 μ m band with a 0.2 cm⁻¹ resolution. Each band spectrum is shown in Fig. 13.1. Polarization data for the two axes are also obtained for each spectral band. The CO₂ absorption bands near 1.6 and 2.0 μ m are quite important because they provide a significant amount of information near the Earth's surface where changes in CO₂ concentrations are most apparent. The 0.76 μ m band is also important to detect ground surface pressure and cirrus cloud information. If satellite observations are to improve over the existing ground network, monthly averaged column data at a precision of 1% or better, for an 8° \times 10° footprint are needed [2]. The GOSAT mission was designed to observe CO₂ density with 1% relative precision averaged in a certain period and with 10 km \times 10 km footprint size during the first commitment period of the Kyoto Protocol (2008–2012).

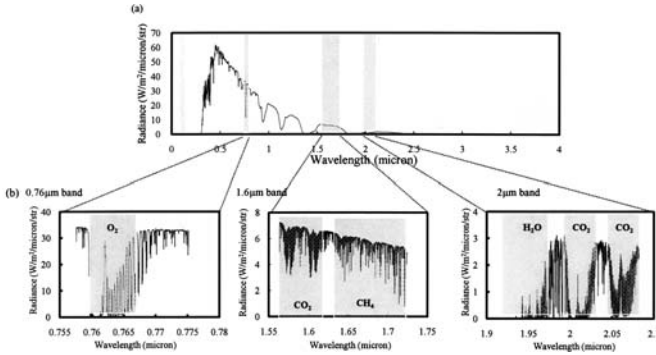


Fig. 13.1 Three-spectral bands of the GOSAT TANSO-FTS-SWIR (b) and solar spectral radiance (a)

NIES developed a GOSAT Data Handling Facility (DHF) for processing GOSAT data in routine operation. After data reception and Level 1 processing by JAXA, GOSAT data will be transferred to the DHF. TANSO-FTS data will provide information for spectral analysis, while TANSO-CAI data will be used to generate cloud and aerosol information. Later, these data will be combined to calculate CO₂ and CH₄ column abundances at observation points with no or only thin clouds and aerosol layers. Furthermore, an atmospheric transport model will be used with the obtained distribution of CO₂ column abundances to estimate global distributions of CO₂ fluxes, as well as to generate three-dimensional distributions of CO₂ concentrations. Retrieved CO₂ column abundances will be distributed to users. It is important for the users to know the precision of the retrieved column abundances. Therefore, it is necessary to clarify the precision of the retrieved CO₂ column abundance.

13.2.3 Previous Error Analysis

In this section, error analysis of the column abundances derived from SCIAMACHY data and IMG data is introduced.

SCIAMACHY [4, 5]

An algorithm to retrieve trace gas vertical columns is first introduced. The Weighting Function Modified Differential Optical Absorption Spectroscopy (WFM-DOAS) is based on fitting the logarithm of a linearized radiative transfer model I_i^{mod} plus a low-order polynomial P_i to the logarithm of the observed sun-normalized radiance I_i^{obs} , which is the ratio of the observed nadir radiance and the solar irradiance spectrum. Index i refers to the detector pixel number i . Provided there exists an appropriate spectra fitting window, I_i^{obs} , depends on the true but unknown vertical columns of

the trace gases of interest (components of vector \mathbf{V}^T). The WFM-DOAS equation can be written as follows:

$$\ln I_i^{obs}(\mathbf{V}^T) - \ln I_i^{mod}(\bar{\mathbf{V}}) + \sum_{j=1}^J \frac{\partial \ln I_i^{mod}}{\partial V_j} |_{\mathbf{V}_j} \times (\hat{\mathbf{V}}_j - \bar{\mathbf{V}}_j) + P_i(a_m) \equiv \| RES_i \|^2 \rightarrow min. \quad (13.1)$$

The fit parameters are the desired trace gas vertical columns \hat{V}^j and the polynomial coefficients a_m . The errors of the retrieved columns have been calculated as follows

$$\sigma_{\hat{V}_j} = \sqrt{(C_x)_{jj} \times \sum_i RES_i^2 / (nv - m)}, \quad (13.2)$$

where $(C_x)_{jj}$ is the j -th diagonal element of the covariance matrix, nv is the number of spectral points in the fitting window, and m is the number of linear fit parameters.

The errors derived from the error of input parameters are examined. A preliminary version of the WFM-DOAS algorithm was implemented based on a fast look-up table approach. The fast look-up table approach introduces quantifiable errors [5], which are related to the (rather sparse) grid selected for the reference spectra look-up table: solar zenith angle interpolation, scan angle correction, and surface elevation (pressure). In addition, factors affecting the calculated model spectrum are surface albedo, surface pressure, aerosols, vertical CO_2 profiles, water vapor, and temperature. When varied alone, each of these parameters creates an error in the column of typically less than 2% [2]. Although the effect of each parameter on the retrieved column abundance is evaluated, the total error on the retrieved column abundance, which includes the error source written above, is not evaluated. Furthermore, a modified retrieval algorithm called the (FSI)-WFM-DOAS was developed. The (FSI)-WFM-DOAS algorithm generates a reference spectrum for every single SCIAMACHY measurement to obtain the best linearization point for retrieval. Analysis of the (FSI)-WFM-DOAS retrievals with respect to the ground-based, FTIR instrumentation (located at Egbert, Canada) reveals that the overall bias of the CO_2 columns retrieved by the FSI algorithm is <4.0% with a monthly precision of 1×1 gridded data close to 1.0% [3].

IMG [14]

The retrieval algorithm uses the maximum likelihood method to take into account the statistical prior knowledge into the least squares method. An iterative calculation is used, so that the estimator $\hat{\mathbf{X}}$ will converge to the global minimum point of the error curve.

$$\mathbf{e}_y^T \mathbf{S}_\epsilon^{-1} \mathbf{e}_y + \mathbf{e}_x^T \mathbf{S}_a^{-1} \mathbf{e}_x, \quad (13.3)$$

$$\mathbf{e}_y = (\mathbf{y} - \mathbf{y}_0) - \mathbf{K}(\mathbf{X} - \mathbf{X}_0), \quad (13.4)$$

$$\mathbf{e}_x = (\mathbf{X} - \mathbf{X}_0), \quad (13.5)$$

where \mathbf{y} , which has as many rows as the number $n\nu$ channel, is a vector of observations (radiance); \mathbf{X}_0 obtained from the analysis of climatological data is used as an initial guess for $\hat{\mathbf{X}}$; \mathbf{y}_0 is a vector of radiance associated with \mathbf{X}_0 ; \mathbf{K} is the Jacobian matrix for the $n\nu$ channels and m estimation parameters; \mathbf{S}_ϵ is the error covariance matrix and includes the observed noise and forward model error; and \mathbf{S}_a is the covariance matrix of \mathbf{X}_0 . In an iterative method, the k -th estimator $\hat{\mathbf{X}}_k$ is used as the linearization point for the next iteration. $\hat{\mathbf{X}}_{k+1}$ is calculated as follows:

$$\hat{\mathbf{X}}_{k+1} - \hat{\mathbf{X}}_k = \mathbf{C}_k(\mathbf{y} - \mathbf{y}_k) + (\mathbf{I} - \mathbf{C}_k\mathbf{K}_k)(\mathbf{X}_0 - \hat{\mathbf{X}}_k) \quad (13.6)$$

$$\mathbf{C}_k = (\mathbf{K}_k^T\mathbf{S}_{\epsilon,k}^{-1}\mathbf{K}_k + \mathbf{S}_a^{-1})^{-1}\mathbf{K}_k^T\mathbf{S}_{\epsilon,k}^{-1}, \quad (13.7)$$

where \mathbf{C}_k , \mathbf{K}_k , and $\mathbf{S}_{\epsilon,k}$ are calculated using $\hat{\mathbf{X}}_k$. Iteration can reduce the nonlinearity effects, and the final estimation is obtained when sufficient convergence is achieved. The covariance of estimation errors \mathbf{S}_x is written as

$$\mathbf{S}_x = (\mathbf{K}^T\mathbf{S}_\epsilon^{-1}\mathbf{K} + \mathbf{S}_a^{-1})^{-1}. \quad (13.8)$$

It should be noted that the evaluation of the retrieved column abundances using SCIAMACHY data and IMG data are performed above. However, input parameters may suffer when column abundances are retrieved. For example, the temperature profile and water vapor amount profile in the atmosphere at the time of spectrum observation affect the intensity of irradiance that is noted by the sensor. Therefore, the temperature profile and water vapor amount should be known precisely in order to retrieve CO_2 column abundances. However, it is difficult to know these input parameters. Therefore, it is better to consider that the retrieved column abundances have an error due to the error of the input parameters. Thus, the precision of the retrieved column abundances is included in this study.

13.3 Error Evaluation and Results

Section 13.3.1 introduces the retrieval algorithm used for this study, which is based on the Rodgers method [20], while Sect. 13.3.2 provides a discussion of the error evaluation method for this retrieval algorithm. Finally, Sect. 13.3.3 provides an example of the observation scenario, such as measurement geometry and the results of error evaluation. It should be noted that this retrieval algorithm may be modified in the future.

13.3.1 Retrieval Method

CO_2 and CH_4 column abundances are retrieved separately. The retrieval method of CH_4 column abundances will be different from that of CO_2 column abundances in terms of the unknown and input parameters. This section focuses on the retrieval

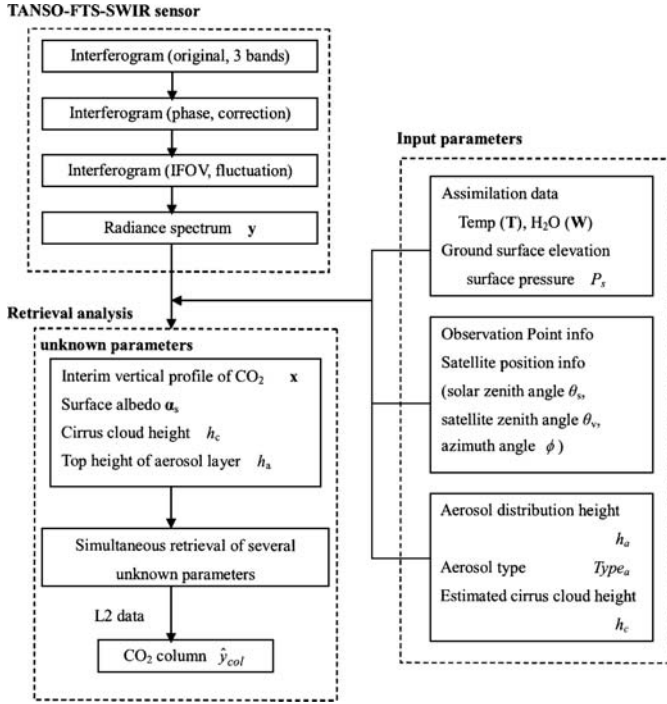


Fig. 13.2 Flow until a CO₂ column abundances is retrieved

of CO₂ column abundance. Figure 13.2 depicts the retrieval test flow until the CO₂ column abundance is retrieved. In this study, the unknown parameters are CO₂ density for each layer \mathbf{x} , albedo of the observation region α_s , cirrus cloud optical depth at $0.55 \mu\text{m}$ τ_c , and aerosol optical depth at $0.55 \mu\text{m}$ τ_a . Clouds were assumed to be thin cirrus. Since albedo varies according to wave number, albedos were estimated at 5 wave number points in the band at even intervals used in the retrieval.

Input parameters are as follows. The radiance spectrum obtained by TANSO-FTS is denoted as \mathbf{y} . Input parameters related to measurement geometry are the solar zenith angle θ_s , the satellite’s zenith angle θ_v , and azimuth angle between the sun and satellite ϕ . Cloud height h_c is a parameter that provides information about cloud conditions. Parameters related to atmospheric conditions are surface pressure P_s , temperature profile \mathbf{T} , and water vapor amount \mathbf{W} . Aerosol parameters are aerosol height distribution (h_a) and aerosol type ($Type_a$). In practice, representing aerosols is difficult due to the different optical characteristics derived from varying aerosol compositions. Therefore, aerosols are expressed as typical rural and urban aerosols. The relationship between unknown and input parameters is represented by introducing the following function f with error \mathbf{e}_X

$$(\mathbf{x}\alpha_s\tau_c\tau_a)^T = f(\mathbf{y}, \theta_s, \theta_v, \phi, h_c, P_s, \mathbf{T}, h_a, \mathbf{W}, Type_a) + \mathbf{e}_X. \quad (13.9)$$

The quantities to be retrieved can be represented by the state vector $\mathbf{X}(=\mathbf{x} \ \alpha_s \ \tau_c \ \tau_a)^T$. \mathbf{y} is approximated by some forward model $F(\mathbf{X})$. The relationship between the radiance spectrum \mathbf{y} and the state vector \mathbf{X} is written with error covariance \mathbf{e}_y ,

$$\mathbf{y} = F(\mathbf{X}) + \mathbf{e}_y. \quad (13.10)$$

A linear problem is one for which the forward model is linear: $\mathbf{y}=\mathbf{K}\mathbf{X}$ where $\mathbf{K}=\partial F(\mathbf{X})/\partial\mathbf{X}$. Gaussian statistics usually provide a good approximation for the errors in real measurement. The maximum posteriori approach is considered for this nonlinear problem. The Bayesian solution for the linear problem can be easily modified for an inverse problem in which the forward model is a general function of the state vector \mathbf{X} . The measurement error \mathbf{e}_y in (13.10) is assumed to be Gaussian, and there is a prior estimate with a Gaussian error

$$-2 \ln P(\mathbf{X}|\mathbf{y}) = [\mathbf{y} - F(\mathbf{X})]^T \mathbf{S}_\epsilon^{-1} [\mathbf{y} - F(\mathbf{X})] + [\mathbf{X} - \mathbf{X}_0]^T \mathbf{S}_a^{-1} [\mathbf{X} - \mathbf{X}_0] + c, \quad (13.11)$$

where $P(q)$ is expressed as a probability distribution function *pdf* of q , and c is constant. As in the linear case, the task in the nonlinear case is to find a best estimate $\hat{\mathbf{X}}$ and an error characteristic that describes *pdf* well enough for practical purposes. The Gauss–Newton method gives

$$\mathbf{X}_{k+1} = \mathbf{X}_k + (\mathbf{S}_a^{-1} \mathbf{K}_k^T \mathbf{S}_\epsilon^{-1} \mathbf{K}_k)^{-1} \{ \mathbf{K}_k^T \mathbf{S}_\epsilon^{-1} [\mathbf{y} - F(\mathbf{X}_k)] - \mathbf{S}_a^{-1} [\mathbf{X}_k - \mathbf{X}_0] \}, \quad (13.12)$$

where k is the iteration number, \mathbf{X}_0 is a prior of the state vector, \mathbf{S}_a is the associated covariance matrix of \mathbf{X}_0 , and \mathbf{S}_ϵ is the error covariance.

nl refers to the number of layers and vector $\hat{\mathbf{x}}$ ($=\hat{x}_1 \hat{x}_2 \dots \hat{x}_{nl}$)^T refers to the retrieved CO₂ density for all layers. CO₂ column abundances \hat{y}_{col} is obtained from $\hat{\mathbf{x}}$ by [10].

$$\hat{y}_{col} = \int_0^{z_{top}} \hat{x} n_a dz, \quad (13.13)$$

where z is referred to the altitude, z_{top} is the altitude at top layer, and n_a is the number of density of air.

13.3.2 Error Evaluation

Since the CO₂ column abundance evaluation method is the same as that for CH₄ column abundance, the term ‘‘column abundance’’ in this section indicates both CO₂ and CH₄ column abundance. The variance of the retrieved column abundances, σ^2 , is estimated, and the error bar is assumed to be provided by 1-sigma analysis, $\sigma(= \sqrt{\sigma^2})$. Internal and external errors are also considered. Internal error is calculated from the residuals for spectral fitting using the Gauss–Newton approach in (13.12). σ_{int}^2 denotes the variance of retrieved column abundances due to internal error. External error, which is the error of the retrieved column abundances due to

the errors in the input parameters, is written as σ_{ext}^2 . The variance of total error σ^2 is defined as the sum of σ_{int}^2 and σ_{ext}^2 . σ^2 is given by

$$\sigma^2 = \sigma_{int}^2 + \sigma_{ext}^2. \quad (13.14)$$

Internal and external errors are calculated in the following section.

Internal Error

Column abundances \hat{y}_{col} are obtained numerically using the trapezoid rule to integrate (13.13) as

$$\hat{y}_{col}(\hat{\mathbf{x}}) = \mathbf{h}^T \hat{\mathbf{x}}, \quad (13.15)$$

where

$$\mathbf{h} = \begin{pmatrix} 1/2n_{a,1} \Delta z_1 \\ 1/2n_{a,2} (\Delta z_1 + \Delta z_2) \\ \vdots \\ 1/2n_{a,nl-1} \Delta z_{nl-1} \end{pmatrix},$$

Δz_l expresses $z(l+1) - z(l)$ for a certain layer l . The variance of the random variable q is written as $Var[q]$, and the variance of $\hat{y}_{col}(\hat{\mathbf{x}})$ is given by (13.15) as

$$Var[\hat{y}_{col}(\hat{\mathbf{x}})] = Var[\mathbf{h}^T \hat{\mathbf{x}}] = \mathbf{h}^T Var[\hat{\mathbf{x}}] \mathbf{h} = \sigma_{int}^2, \quad (13.16)$$

where $Var[\hat{\mathbf{x}}]$ is given as

$$Var(\hat{\mathbf{x}}) = (\mathbf{K}^T \Sigma^{-1} \mathbf{K} + \mathbf{S}_a^{-1})^{-1}. \quad (13.17)$$

It is assumed that the elements of \mathbf{y} are independent of each other, and Σ is given by $\Sigma = \psi \mathbf{S}_\epsilon$. ψ is written as

$$\psi = \frac{(\mathbf{y} - F(\hat{\mathbf{X}})) \mathbf{S}_\epsilon^{-1} (\mathbf{y} - F(\hat{\mathbf{X}}))}{nv - m}, \quad (13.18)$$

where nv is the number of channels in \mathbf{y} when \mathbf{X} is retrieved, and m is the number of unknown parameters. Consequently, σ_{int}^2 can be obtained by

$$\sigma_{int}^2 = \mathbf{h}^T \left(\frac{\mathbf{K}^T \mathbf{S}_\epsilon^{-1} \mathbf{K}}{\psi} + \mathbf{S}_a^{-1} \right)^{-1} \mathbf{h}. \quad (13.19)$$

External Error

Next, the effects on retrieved CO₂ column abundances derived from the errors of input parameters are considered. External error was not reflected in the error evaluations of the retrieved CO₂ column abundances derived from SCIAMACHY

and IMG. The levels of all factors in the experiment represent only a random subset of the possible factor levels of interest [15]. Randomization is a procedure whereby factor-level combinations are assigned to a test sequence in such a way that every factor-level combination has an equal chance of being assigned to any experiment unit or position in the test sequence. In a completely randomized design, all the factor-level combinations in the experiment are randomly assigned to experimental units. However, complete factorial experiments cannot always be conducted due to economic, time, and other constraints. This paper makes use of an orthogonal array to obtain fractional factorial experiments, since using orthogonal arrays is a widely-accepted method used in industry, with σ_{ext}^2 obtained from factorial experiments using an orthogonal array. The model terms that represent random factor levels are themselves random variables. When error factors are denoted by A, B, \dots , the model can be expressed as follows:

$$\hat{y}_{col} = \mu(A, B, \dots) + e_{col} = \mu + \alpha + \beta + \dots + e_{col}, \quad (13.20)$$

where e_{col} ($\sim N(0, \sigma_e^2)$) represents random error effects, α, β, \dots represent the contribution to \hat{y}_{col} derived from factors A, B, \dots , respectively, and " $\sim N(0, \sigma_e^2)$ " indicates that the function has a normal distribution with mean 0 and variance σ_e^2 . All of the random variables for A, B, \dots are assumed to be statistically independent. When the factor effects are all random, the response \hat{y}_{col} has a common normal distribution with mean $\mu(A, B, \dots)$ and variance σ_{ext}^2

$$\hat{y}_{col} \sim N(\mu(A, B, \dots), \sigma_{ext}^2). \quad (13.21)$$

σ_{ext}^2 is obtained from the experimental results based on an orthogonal array. The number of tests is represented by N , and σ_{ext}^2 is obtained by

$$\sigma_{ext}^2 = \frac{1}{N} \sum_{n=1}^N (\hat{y}_{col,n} - \bar{y}_{col})^2, \quad (13.22)$$

where \bar{y}_{col} is the mean of retrieved column abundances from N tests. That is,

$$\bar{y}_{col} = \frac{1}{N} \sum_{n=1}^N \hat{y}_{col,n}. \quad (13.23)$$

13.3.3 Error Evaluation Results

Setting Parameters

Table 13.2 shows the measurement geometry and atmospheric conditions. The atmosphere is represented using the US Standard model. It was assumed that there are

Table 13.2 Measurement geometry and atmospheric condition during observation

	Parameter	Value
(a) Observation geometry	Solar zenith angle	30.0°
	Satellite zenith angle	0° (nadir observation)
	Atmospheric model	US Standard model
	CO ₂ density	380 ppmv for all layers
(b) Atmospheric condition	Cirrus height	10 km
	Aerosol type	Rural type aerosol
	Aerosol optical depth	0.1

thin cirrus clouds with 0.2 optical depth at 0.55 μm and at a middle height of 10 km, and that aerosols are distributed uniformly from the Earth's surface to an altitude of 3 km with 0.1 optical depth at 0.55 μm . CO₂ density is low at the higher altitude, and it is difficult to retrieve CO₂ density at such altitudes precisely. Therefore, the number of layers nl is set to 15 from the Earth's surface to 15 km at 1-km intervals. It is assumed that the observation area on the Earth's surface is covered with conifer forests. These parameters are given as true values of the estimation parameters when observation spectra are calculated by a radiative transfer code.

CO₂ column abundances are retrieved from the 1.6 μm band only to reduce the amount of time required. The 1.6 μm GOSAT spectrum y is calculated in the following manner. Solar incident radiance every 0.01 cm^{-1} from 6,180 to 6,380 cm^{-1} is calculated using the HSTAR (High resolution RSTAR [17, 18]) code, which calculates the spectral radiance of a spectral band using the line-by-line method according to cloud and aerosol conditions. Polarization bands are not used here, and solar radiance multispectral scattering in the atmosphere is taken into consideration. An optical filter function of the 1.6 μm band of the sensor is applied to the incident radiance data. Interfered radiance, called an interferogram, is obtained from the FTS by calculating an inverse Fourier transformation. The spectrum is obtained every 0.2 cm^{-1} using a Fourier transformation. Then, sensor noise corresponding to $S/N = 300$ is added to the obtained spectrum, since the sensor efficiency is $S/N = 300$. It is assumed that the sensor characteristic is known based on an examination before the satellite is launched. Essentially, this yields a maximum spectrum y_{max} and standard deviation of noise σ_ϵ obtained by $y_{max}/300$. Noise is assumed to follow a normal distribution with mean value 0 and variance σ_ϵ^2 , and is calculated for every 0.2 cm^{-1} of the spectrum.

\mathbf{S}_a and \mathbf{S}_ϵ in (13.12) is set as follows. Knowing \mathbf{S}_a correctly is difficult in practice. In the case where the elements of \mathbf{S}_a are small, retrieval precision greatly depends on \mathbf{S}_a . Therefore, this retrieval algorithm uses a larger \mathbf{S}_a than that which is normally practical. \mathbf{S}_a variance terms are shown in Table 13.3. Covariance terms of \mathbf{S}_a are set to 0, and it is assumed that \mathbf{S}_ϵ is derived only from sensor efficiency. Then, all \mathbf{S}_ϵ variance terms are set to σ_ϵ^2 , and \mathbf{S}_ϵ covariance terms are set to 0.

The number of factors in the experiment to obtain σ_{ext}^2 is set as follows. The nine input parameters of (13.9) are candidate factors in the experiment. The direction of the sun is calculated using either an "Astronomical Almanac [19]" or "Japanese

Table 13.3 S_a variance terms

		x					(ppmv ²)	
Layer	1	2	3	4	5	6	7	8
Value	97.31 ²	86.79 ²	77.41 ²	69.04 ²	61.57 ²	54.92 ²	48.98 ²	43.68 ²
Layer	9	10	11	12	13	14	15	
Value	38.96 ²	34.75 ²	30.99 ²	27.64 ²	24.65 ²	21.99 ²	19.61 ²	
		α_s					τ_c	τ_a
No.	1	2	3	4	5	1	1	
Value	1 ²	1 ²	1 ²	1 ²	1 ²	0.2 ²	0.2 ²	

Ephemeris [11]” with high precision. Satellite positions are also obtained quite accurately from orbit information acquired by a global positioning system. Consequently, errors of θ_s , θ_v , and ϕ are not considered in this study, leaving six input parameters, which, except for θ_s , θ_v , and ϕ , are used as error factors. Table 13.4 shows the levels for each set error factor. Levels for each error factor are set as follows. Levels for each factor are set to the accuracies of each input parameter. Input parameters of temperature profile **T**, water vapor profile **W**, and surface pressure P_s are used as provided by six hourly ($1.25^\circ \times 1.25^\circ$) Global Analysis data (GANAL) from the Japan Meteorological Agency. It is assumed that the errors of GANAL data for **T** and **W** are fewer than 2 K and 30% of **W**, respectively. P_s precision is assumed to be 3 hPa, and P_s error is also assumed to be the error of the estimated surface elevation of the observation area. “Temperature profile shift” levels are set to 0 K (level 1), -2 K (level 2), and $+2$ K (level 3). CO₂ column abundance retrievals for each level are conducted by adding levels to the temperature profile of the GANAL data for all layers. “Water vapor amount shift” levels are set to 0% (level 1), -30% (level 2), and $+30\%$ (level 3) of the GANAL data. Retrievals for each level are carried out by adding levels to the water vapor amount of the GANAL data similar to the method for the “temperature profile shift”. Atmospheric pressures for each layer from the surface to the top of the atmosphere are changed according to hydrostatic equilibrium using information on temperature shift and water vapor shift for each level. Furthermore, the shifts of atmospheric pressure at the surface are set to 0 hPa (level 1), -3 hPa (level 2), and $+3$ hPa (level 3). Cases where aerosol height distributions differ from the practical distribution when the spectrum was obtained were also considered with “aerosol height distribution” levels uniformly set to 3 km from the surface (level 1), 2 km from the surface (level 2), and 4 km from the surface (level 3). GOSAT is designed to estimate cirrus cloud height from the A-band of oxygen and saturated absorption region of water vapor in the 2.0 μm band before CO₂ column abundances are retrieved. Cloud top pressure changes based on cloud height, and it is assumed the precision of the estimated cirrus height is ± 1 km. “Shift of cirrus cloud height” levels are set to 0 km (level 0), -1 km (level 2), and $+1$ km (level 3). Experiments for each level are conducted by adding the level to the cloud height that is regarded as the estimated cirrus height. For the factors given above, there are

Table 13.4 Trace gas observation sensors onboard satellite

Label	Factor		Level		
			1	2	3
<i>A</i>	Cirrus height (km)	h_c	0	-1	+1
<i>B</i>	Surface pressure (hPa)	P_S	+0	-3	+3
<i>C</i>	Temperature profile shift (K)	T	0	-2	+2
<i>D</i>	Aerosol height distribution (km)	h_a	0-3	0-2	0-4
<i>E</i>	Water vapor amount shift (%)	W	0	-30	+30
<i>F</i>	Aerosol type	$Type_a$	Rural	Urban	

Table 13.5 Orthogonal array L18

Text no.	Level						Text no.	Level					
	<i>A</i>	<i>B</i>	<i>C</i>	<i>D</i>	<i>E</i>	<i>F</i>		<i>A</i>	<i>B</i>	<i>C</i>	<i>D</i>	<i>E</i>	<i>F</i>
1	1	1	1	1	1	1	10	1	1	3	3	2	2
2	1	2	2	2	2	1	11	1	2	1	1	3	2
3	1	3	3	3	3	1	12	1	3	2	2	1	2
4	2	1	1	2	2	1	13	2	1	2	3	1	2
5	2	2	2	3	3	1	14	2	2	3	1	2	2
6	2	3	3	1	1	1	15	2	3	1	2	3	2
7	3	1	2	1	3	1	16	3	1	3	2	3	2
8	3	2	3	2	1	1	17	3	2	1	3	1	2
9	3	3	1	3	2	1	18	3	3	2	1	2	2

Table 13.6 Initial values of unknown parameters

Unknown parameter	Value
CO ₂ density profile	370 ppmv for all layers
Surface albedo	0.3 for all spectral points
Cirrus optical depth	0.05
Aerosol optical depth	0.1

three levels. Since the surface of the observation area is covered by conifer forest, “aerosol type” levels are set to rural type (level 0) and urban type (level 1), giving a total number of two possible levels. A major difference between “rural” and “urban” aerosols is the amount of black carbon particles. Table 13.5, “L18 (Latin square 18) orthogonal array” presents the orthogonal array used to obtain σ_{ext}^2 . The number of tests N is 18. Initial values of estimation parameters are presented in Table 13.6. CO₂ column abundances are retrieved using (13.12). Since the solar Fraunhofer line mask and the water vapor mask are applied to 0.2 cm⁻¹ resolution spectrum, the number of channels in y is 232 when CO₂ column abundances are retrieved. The iteration is conducted until the change of ψ is less than 10⁻⁵.

Results

The evaluation results based on the input parameters are shown below. It is assumed that **T**, **W**, and P_s of the GANAL data are obtained to the same accuracy as the true values (US Standard values): the aerosols are distributed from the surface to 3 km; the estimated cloud height is 10 km; and the true CO₂ column abundance, which is not known in practical observation, is 7.109×10^{21} mol · cm⁻². Retrieved CO₂ column abundance is 7.118×10^{21} mol · cm⁻², and the variance of the internal error, σ_{int}^2 , using (13.16) is 4.451×10^{38} mol · cm⁻². The variance of the external error is obtained by experiments following the orthogonal design table in Table 13.5. Retrieved CO₂ column abundances \hat{y}_{col} for each test are shown in Fig. 13.3. A mean value of \hat{y}_{col} from 18 tests is 7.135×10^{21} mol · cm⁻², and σ_{ext}^2 is 1.133×10^{38} mol · cm⁻². The variance of total error is 5.584×10^{38} mol · cm⁻², with an error bar of 2.363×10^{19} mol · cm⁻². The CO₂ column abundance is $7.118 \times 10^{21} \pm 0.024 \times 10^{21}$ mol · cm⁻².

True CO₂ column abundances are covered by the error bars. The error ratio, which is calculated by $\sigma \times 100 / \hat{y}_{col}$, is 0.33%. It is conceivable that a precision of less than 1% error was achieved for the retrieved CO₂ column abundance in this observation scenario. Since σ_{ext}^2 , which is obtained by experiments using orthogonal arrays, takes a long time to calculate, σ_{ext}^2 will be obtained in advance. Therefore, it is necessary to obtain σ_{ext}^2 under several scenarios. GOSAT is designed to retrieve CO₂ column abundance data by both nadir observation and observations in the cross-track direction, so error evaluation needs to be conducted by altering solar zenith angle, satellite zenith angle, and azimuth angle between the sun and satellite using numerical simulation. Furthermore, in addition to rural and urban type aerosols, there are other types of aerosols, such as yellow sand and sea spray type aerosols. Therefore, it is necessary that retrieval precision is evaluated for different aerosol conditions. GOSAT will also take spectrum measurements using sun glint from large bodies of water.

Since the retrieval algorithm to be used in the DHF is currently being developed and coded, it is important to know the contribution of input parameter precision

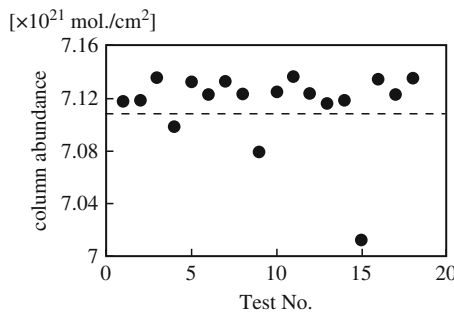


Fig. 13.3 Retrieved CO₂ column abundances for each test following orthogonal array

Table 13.7 Initial values of unknown parameters

Level	Factor	$S \times 10^{39}$	η	$V \times 10^{39}$	F -value	ρ (%)
<i>A</i>	Cirrus height	0.019	2	0.010	0.11	0.952
<i>B</i>	Surface pressure	0.119	2	0.059	0.70	5.816
<i>C</i>	Temperature profile shift	0.559	2	0.279	3.30	27.393
<i>D</i>	Aerosol height distribution	0.213	2	0.107	1.26	10.449
<i>E</i>	Water vapor amount shift	0.397	2	0.198	2.34	19.448
<i>F</i>	Aerosol type	0.224	1	0.224	2.65	11.003
	Error	0.509	6	0.085		24.938
	Total	2.039	17	0.120		

to the retrieved CO_2 column abundances to improve the retrieval precision, that is, it was assumed that the retrieval precision is improved by reducing the largest factor's contribution to the retrieved CO_2 column abundances. Next, the magnitude of contribution to the retrieved CO_2 column abundance for each factor is calculated derived based on an analysis of variance (ANOVA) using retrieved CO_2 column abundances shown in Fig. 13.3, and the results are presented in Table 13.7. S , η , V , F -value, and ρ in Table 13.7 stand for sums of squares, degrees of freedom, mean squares, F -value, and the rate of contribution (%), respectively. The sum of squares for factor "Total" S_{Total} is calculated by (13.22) $\times N$. When Q represents factors A , B , C , D , and E , which have three levels, the sums of squares S_Q are

$$S_Q = 6 \sum_{q=1}^3 (\hat{y}_{col,q} - \bar{y}_{col})^2, \quad (13.24)$$

where q refers to the level, and $\bar{y}_{col,q}$ is the mean value of \hat{y}_{col} when the level of factor " Q " is q . When Q represents the factor F , which has two levels,

$$S_Q = 9 \sum_{q=1}^2 (\bar{y}_{col,q} - \bar{y}_{col})^2. \quad (13.25)$$

Furthermore, the "Error" factor of sum of squares S_{Error} is obtained by

$$S_{Error} = S_{Total} - S_A - S_B - S_C - S_D - S_E - S_F. \quad (13.26)$$

and V , F -value, and ρ of factor q are obtained by S_q/η , V_q/V_{Error} , and S_q/S_{Total} , respectively.

Based on Table 13.7, "Temperature profile shift" is the largest contributor, followed by "water vapor amount shift". On the other hand, "cirrus height" and "surface pressure" are low contributors when the estimated cirrus height error and the surface pressure shift are given within ± 1 km and ± 3 hPa. Therefore, these two factors are not included in the error, and the number of factors can be reduced. Thus, it can be concluded that ANOVA effectively examines the contribution of each error

term to the retrieved CO₂ column abundances. Prior to launch of the satellite, it is necessary to conclusively determine the contribution of each of the error factors on the retrieved CO₂ column.

13.4 Conclusions

The Greenhouse Gases Observing Satellite (GOSAT), which has been slated to be launched in early 2009, will be the world's first satellite to collect data on the concentrations of carbon dioxide (CO₂) and methane (CH₄), two major greenhouse gases, from space. GOSAT features an onboard down-looking FTS sensor. The data collected will be distributed to users in the form of digital data and global maps of CO₂ and CH₄ column abundances.

Clarifying the precision of the retrieved CO₂ and CH₄ column abundances for GOSAT users is an important component in developing an understanding of global warming mechanisms. This paper describes in detail an evaluation method to clarify GOSAT data precision by setting a measurement scenario and performing a numerical simulation to judge the precision of retrieved CO₂ column abundance data. The evaluation method presented considers two types of error: internal error derived from spectral fitting residuals, and external error derived from the error of input parameters when the column abundances are retrieved. Internal errors were calculated analytically, while the external errors, which are not considered in the error estimation using IMG data and SCIAMACHY data, were obtained using factorial experiments.

The following observation scenario provided the basis for this analysis. Cirrus clouds and aerosols were present in the atmosphere and CO₂ column abundance data was retrieved using nadir observation. The spectra calculated were assumed to be GOSAT-measured spectra. Eighteen experiments were conducted by changing combinations of levels according to an orthogonal array, yielding 18 CO₂ column abundances obtained using numerical simulation. The precision of the CO₂ column abundances was evaluated, and the results were shown. Based on the above scenario, less than 1% precision was demonstrated for the retrieved CO₂ column abundances. Analysis of variance was applied to determine the magnitude of the contribution to the retrieved CO₂ column abundances. In the future, the magnitude of contribution will be examined for several measurement scenarios.

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