

Plant traits and population characteristics predict extinctions in a long-term survey of Mediterranean annual plants

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Abstract Global and local environmental changes lead to frequent plant extinctions many of which occur in man-made habitats such as agricultural fields. Plant traits and site conditions modify risks of extinction, but strength and sense of their effect are not known yet. Here, we present a long-term survey of population sizes for Mediterranean annual plants that we revisited 20 years after their first record to evaluate climate, population size, traits and habitat requirements as drivers of local extinctions. Small populations had an increased probability of extinction in our data-set. Our analyses revealed that seed production and survival of seeds in the soil seed bank decreased extinction rate, whereas plant size increased extinction probability. Mean annual temperature increased extinction rates of annual plants in cereal fields. We discuss these effects as a response to recent and ongoing habitat changes, and discuss how traits may be used to guide conservation practices in the face of local extinctions.

Keywords Plant size · Seed banks · Local extinction · Seed number · Specific leaf area

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Introduction

Global environmental changes such as climate warming, changing land use and habitat fragmentation are identified as major reasons for the decline of plant diversity (Bobbink et al. 1998; Rogers et al. 2008; Römermann et al. 2008; Van Calster et al. 2008; Wiegmann and Waller 2006). However, plants differ in their specific extinction risks according to their functional traits (Farnsworth and Ogurcak 2008; Fréville et al. 2007; Kahmen et al. 2002; Lavergne et al. 2006; Ozinga et al. 2008; Piqueray et al. 2011; Römermann et al. 2008). Moreover, works on plant communities showed how traits interact with environment to trigger different extinction risks among plant species (Fréville et al. 2007; Kahmen et al. 2002; Römermann et al. 2009). Traits identified to trigger major population biological functions include specific leaf area (SLA), plant height, seed size and flowering date, interacting with management, habitat size and nutrient level (Fréville et al. 2007; Kahmen et al. 2002; Römermann et al. 2009). Works on regional scales suggest that relevant traits to explain interspecific differences in extinction risk depend on specific environmental conditions (Fréville et al. 2007; Lavergne et al. 2006; Ozinga et al. 2008; Rogers et al. 2008; Römermann et al. 2008; Wiegmann and Waller 2006). However direction and strength of effects of functional traits on extinction are not well known.

Indeed, relevant effects of traits might be detected when comparing local habitat and population properties, such as population size and mating system, which both have strong effects on population genetic structure (Duminil et al. 2007; Ellstrand 1992). Moreover, the identification of intrinsic risk factors (traits) for extinction in plants also enables to identify the relevant processes that determine local extinction risk. Therefore, one way to generalize results of extinction risks linked to declining populations across sites is to study specifically functional traits together with population sizes and environmental conditions.

Environmental conditions can change in an unidirectional way driving species to extinction, e.g., through changed disturbance regimes in anthropogenic habitats such as arable fields (Fried et al. 2009a). Environment also show temporal variability that may enhance coexistence of populations via storage effect which is linked to specific plant traits (Facelli et al. 2005; Levine and Rees 2004). Both processes can concern the same factor (e.g., soil moisture) and may be on the same temporal scale (years). For example, soil moisture can directly decrease seed populations in the soil via increasing fungal attack (Blaney and Kotanen 2001; Saatkamp et al. 2014; Schafer and Kotanen 2003). Other factors such as higher nutrient availability and warming of climate have also been shown to change plant community composition and diversity (Bobbink et al. 1998). Thus, directional changes in soil moisture, temperature and nutrient status are interacting with plant traits to influence extinction dynamics of populations for annual plants (Fréville et al. 2007).

Population characteristics known to correlate with extinction risk are population size, genetic structure and diversity. For example, population size influences local extinction, with small populations going extinct faster than large ones (Matthies et al. 2004). Since this makes it difficult to disentangle which factors finally drive a local plant population to extinction, we suggest in this study to use both, plant traits as well as environmental factors and population size in order to go further in the understanding of local plant extinctions. Plant traits can directly modify population dynamics (Angert et al. 2007; Gardarin et al. 2010), for example, adaptations at the seed level and soil seed mortality changes population size of annual plants (Adams et al. 2005; Kalisz and McPeck 1993; Menges 2000; Schmid and Matthies 1994). This extinction risk can be related to their short life cycle, differences in survival in the soil seed bank and to anthropogenic disturbances (Saatkamp

et al. 2011a, 2014). This also separates seed mortality from seed number, in a trade-off with seed size, and hence separates mortality sources in the seed life stage from those in other life stages (Leishman et al. 2000; Moles et al. 2004; Saatkamp et al. 2009). Finally, the rarity of a plant species at larger spatial scales influences its extinction risk, since colonization events strongly depend on abundance of species in the seed rain arriving from larger distances (Hubbell 2001). However, until now, soil seed mortality and seed production have not been clearly separated in their effects on local extinction risk in annual plants (Saatkamp et al. 2009).

Differences in competitive ability among plants as estimated by plant size (Fayolle et al. 2009) or differences in recovery after disturbances related to relative growth rate tightly linked to specific leaf area (Shipley 2006), may help to further understand which processes lead to increased local extinction risk. Finally, also population genetic structure is related to plant traits (Duminil et al. 2007; Ellstrand 1992) and traits might affect extinctions in this way. For instance, the pollen:ovule (P/O) ratio is an indicator trait that relates to gene exchange via breeding system and realized pollination events (Cruden 1977; Loveless and Hamrick 1984), for which we showed elsewhere a link to other risk factors such as land use changes (Lhotte et al. 2014). The P/O ratio also retraces a gradient from low pollen exchanging and inbreeding to high pollen exchanging and obligate outcrossing species (Cruden 1977; Loveless and Hamrick 1984). In the context of rapidly changing habitat conditions, pollination can be a key factor for plants to adapt rapidly and hence we hypothesize that the P/O ratio is in this way a factor that influences independently the risk of plant population extinctions.

Annual plants make up one third of plant diversity in the European Mediterranean region and are at comparatively high risks of extinction (Médail and Verlaque 1997). This extinction risk can be related to their short life cycle, important differences in survival in the soil seed bank and to their habitats typically with strong anthropogenic disturbances (Saatkamp et al. 2011a, 2014). Despite its importance in species number, we know strikingly little about the specific processes for the decline of annual plants in the Mediterranean region. Works that evaluated the relation between traits and local population dynamics or extinctions, did so for a single species (Adams et al. 2005; Kalisz and McPeck 1993). It is therefore expected that comparative analyses of population dynamics for several species will give new insights (Menges 2000). For annual plants of cereal fields, population extinctions are due to changing agricultural practices such as herbicide and fertilizer application as well as higher densities of crop plants (Fried et al. 2009b; Schneider et al. 1994). Annual plants react faster to environmental changes compared to perennial plants and thus effects are easier to detect. Therefore, annual plants in cereal fields give a good opportunity to test the effects of life history traits and habitat requirements on population dynamics.

In this study, we used data on populations for 24 rare and common arable plants together with novel and detailed experimental data on plant traits and environmental parameters to assess local extinction risk in plants. Our approach was to quantify at the same time influence of environmental factors, such as mean annual temperature, population size at the growing sites and characteristics at the species level such as plant traits and habitat requirements.

More specifically, we wanted to answer (i) how plant traits (especially seed production, survival in soil seed bank, seed mass, plant size, SLA and P/O ratio) and population size influence local extinction risk at the long run; (ii) how site climate influences extinction risks; (iii) how water, nutrient, light and temperature requirements as measured by Ellenberg's indicator values influence extinction and (iv) to finally analyse if plants that are rare on regional scales have a higher local extinction risk.

Materials and methods

Study site and species

We gathered data on rare and common cereal weeds in an area of ca. 2500 km² around the Luberon ridge in South Eastern France. Traditional agriculture in this area maintained a high diversity of rare arable plants elsewhere extinct in Europe and data have been gathered in the mid-1980s during field surveys (Filosa 1997). Based on surveys from 1983 to 1985, we revisited populations for 24 species (Supplementary Fig. 1) of rare arable plants. We also used data on rarity levels from a regional flora (Jauzein 1995) to categorise species into either rare or common (Table 1). Nomenclature followed Jauzein (1995).

Climatic variables

Since sites were scattered over large areas without nearby weather stations, we used the worldclim data set together with DIVA-GIS software (Hijmans et al. 2005) to extract climatic variables for our study sites. Study sites were spaced by more than 1 km in most

Table 1 Species studied, plant family, rarity class (Jauzein 1995), IUCN red-list classification for South-Eastern France (Noble et al. 2015): *LC* least concern, *VU* vulnerable, *NT* near threatened and number of populations studied

Species	Family	Rarity	IUCN	Populations
<i>Adonis annua</i>	Ranunculaceae	R	LC	21
<i>Adonis flammea</i>	Ranunculaceae	C	LC	33
<i>Agrostemma githago</i>	Caryophyllaceae	R	LC	9
<i>Androsace maxima</i>	Primulaceae	R	LC	7
<i>Bifora radians</i>	Apiaceae	C	LC	17
<i>Bifora testiculata</i>	Apiaceae	R	NT	5
<i>Bupleurum rotundifolium</i>	Apiaceae	C	LC	3
<i>Camelina microcarpa</i>	Brassicaceae	R	LC	2
<i>Centaurea cyanus</i>	Asteraceae	R	LC	16
<i>Cnicus benedictus</i>	Asteraceae	R	LC	17
<i>Conringia orientalis</i>	Brassicaceae	R	NT	16
<i>Consolida regalis</i>	Ranunculaceae	C	LC	1
<i>Hypocoum pendulum</i>	Papaveraceae	R	VU	3
<i>Legousia hybrida</i>	Campanulaceae	R	LC	5
<i>Legousia speculum-veneris</i>	Campanulaceae	C	LC	4
<i>Myagrum perfoliatum</i>	Brassicaceae	R	LC	2
<i>Neslia apiculata</i>	Brassicaceae	C	LC	32
<i>Nigella damascena</i>	Ranunculaceae	C	LC	1
<i>Papaver argemone</i>	Papaveraceae	R	LC	14
<i>Papaver hybridum</i>	Papaveraceae	R	LC	13
<i>Ranunculus falcatus</i>	Ranunculaceae	R	NT	19
<i>Roemeria hybrida</i>	Papaveraceae	R	VU	10
<i>Turgenia latifolia</i>	Apiaceae	R	NT	28
<i>Vaccaria hispanica</i>	Caryophyllaceae	R	NT	12

cases, translating differences in climate into variance in the data-set. Our study region is characterised by Mediterranean climate with mean annual rainfall varying between 596 and 636 mm, with maxima in April and October. We reduced the manifold climatic variables to mean annual temperature (MAT), since this variable reflects the main gradient in a principal component analysis (PCA) using all climatic variables, and which opposed lowland and upland sites. The first axis of the PCA explained 78% of variance, with MAT and all temperature variables being positively correlated to the first axis and annual precipitation and precipitation variables negatively related except precipitation seasonality; MAT and annual precipitation were strongly negatively correlated ($R^2=0.99$, $p<0.001$). MAT varied from 8 °C in upland to 13 °C in lowland sites.

Population size and local extinction

We used a census data set on rare cereal weeds dating from 1983 to 1985 to gather population sizes and localisations (Filosa 1989). For our analyses we pooled data from 1983 to 1985. In 2005 and 2006 we revisited populations of 24 different plant species. For 290 populations, their size in terms of numbers of individuals has been documented in detail in 1983/1985. These old population sizes enabled us to study its effect on extinction: 192 went extinct and 98 remained stable. The populations were documented by one visit per year at the flowering to fruiting time of the species before wheat harvest. This corresponds to the visits of the initial data collection (D. Filosa, pers. comm.). The census was conducted at the same level of precision as in 1983. For comparison with the 1983 and 1985 data, we also pooled data from 2005 to 2006. In all analyses, we used presence/absence of populations in 2005/2006 as dependent variable. Evidently, the absence of a population in one single year does not equal extinction, since it might be still present in the soil seed bank and this for years (Saatkamp et al. 2009). However, we think that pooling the data over two years of inventory limits the amount of unobserved but still present populations, and that absence of populations in two consecutive years is a reasonable indicator of potential local extinction.

Pollen:Ovule ratio

In spring and summer 2006, we collected three stamens per flower bud, and one bud for each of five randomly chosen individuals for all 24 plant species (illustrated in Supplementary Fig. 1). We stored the three stamens for each bud dry together in Eppendorf tubes. The remaining flower was conserved in 70% alcohol in plastic tubes until we counted number of stamens and ovules. Once all species sampled, we dissolved stamen tissue using 550 μ l of sulphuric acid per tube for 24–48 h. After that, we crushed the remaining tissues with a small glass pestle, added 1650 μ l of water with 2% triton tenside and mixed it. We then centrifuged for 5 min and decanted the superfluous liquid. We added 1000 μ l of alcohol (95%) and mixed it again. We centrifuged again for 5 min, removed liquid, and dried to obtain dry pollen samples. We then added 40 μ l of a 30% sucrose/20% glycerol solution ('counting solution') and sonicated the solution. We counted pollen grains under a microscope using a 1 μ l hematocytometer. We counted ovules under a binocular microscope. We then calculated the number of pollen grains for one stamen as follows: number of pollen grains in 1 μ l \times 40 μ l of counting solution, divided by the three stamens used. We then calculated pollen:ovule ratio (P/O) as the number of pollen grains for one stamen \times number of stamens per flower, divided by the total number of ovules (Cruden 1977).

Seed production, seed mass and plant size

We counted ripe seeds produced for 10 individual plants per species in the field. For species with multi-seeded fruits or infructescences, we counted the number of fruits or infructescences and sampled two of them per individual to count number of seeds per fruit/infructescence. Seed production per individual was then calculated as mean seed number per fruit/infructescences multiplied by the number of fruits/infructescences counted per individual. In our analyses, we used mean seed production for ten individuals as specific trait value. To measure seed mass for each species, we counted three replicates of 10 seeds and weighed them with an analytical balance (± 0.001 mg). At the time of fruiting, we measured plant size (in mm) in the field for 25 individuals chosen randomly in the population, to calculate mean plant size per species.

Soil seed mortality

We studied soil seed mortality using a burial experiment presented in more detail in Saatkamp et al. (2009). For each species, we collected in the field seed material between June and September 2005. We mixed the ripe seeds from at least 10 individuals and stored them dry in paper bags until burial in October 2005. After burial of seeds in nylon mesh bags in a randomised block design (five block replicates), we retrieved buried seeds after 6, 12, 18, 24 and 30 months of burial from October 2005 to April 2008. We determined the viable fraction for every seed lot including an initial value before burial, by using germination tests in a growth chamber and tetrazolium test for remaining seeds (I.S.T.A 1996). This burial experiment yielded mean mortality percentages for a 2.5 year period.

Specific leaf area (SLA)

Leaf material was collected in field on living individuals at begin of flowering (April to June 2007 and 2008) using only green, intact and undamaged leaves. For each species, five individual plants were chosen at random. Two leaves were collected per plant. Until measurement, we put leaves in sealed plastic bags stored at 4 °C, and processed them within 24 h after collection according to the recommendations in Garnier et al. (2001). We measured leaf area only on living and fully turgescient leaves. We used scanned images along with reference surfaces of known size to detect leaf area using the lafore-software package (Lehsten 2005). We then placed leaves in tagged paper bags and dried them at 50 °C for 3 days. We measured dry leaf weight using a fine balance (± 0.001 mg). SLA was calculated for each single leaf as the ratio of fresh leaf area on leaf dry mass. We then used the mean SLA values as plant trait in our analyses.

Ellenberg indicator values

Ellenberg et al. (1992) provided position of a species in gradients of light (L), temperature (T), moisture (F) and nutrients (N) for the central European flora. Indicator values are a very sensible way to characterize habitat conditions for plants (Diekmann 2003; Saatkamp et al. 2011b). We complemented indicator values for *Nigella damascena*, using information

from local floras (Jauzein 1995; Molinier 1981). All trait data are presented in the Supplementary Table 1.

Statistical analyses

Since our data was organized according to three levels of aggregation (populations, sites and species), we run three separate analyses for each level of aggregation i.e.: (1) population size, (2) mean annual temperature aggregated at the site level and (3) plant traits as well as Ellenberg indicator values aggregated at the species level. We first applied a generalized linear regression (GLM) analysis on the three levels of aggregation—populations, sites and species—separately. For the first two levels, we used extinction (1) versus persistence (0) of each population as a binary response variable in a GLM to evaluate effects of population size and site climate and functional traits. All explanatory variables were standardised prior to analyses.

For the species level we used a binomial GLM with logit link (logistic regression) with number of persistence and extinction as success/failure response variable. At the species level we had 24 species and 11 potential explanatory variables, part of which were correlated. We therefore reduced explanatory variables to three use a PCA in the following way. For each of three first PCA-axes with proportions explained variance of 0.25, 0.22 and 0.16, we selected the variable with the highest loading. This retained plant size (PS, loading -0.51 on PCA1), seed number (SN, loading 0.54 on PCA2) and seed bank survival (SB, loading -0.55 on PCA3). Explanatory variables were not correlated according to Pearson's correlation coefficient ($r < 0.5$). We then fitted GLM using the `glm` module in R. The use of species as independent data points is controversial since closely related species show similar features because of common ancestry. In our data set, where several major phylogenetic lineages are represented by only a few representatives, and with many closely related species pairs (Table 1) with contrasting traits, we think it is not reasonable to expect phylogenetic 'bias' in the data, which would further be difficult to quantify with such a small set of species. Therefore we did not perform an analysis including species groupings as random effects.

In supplementary document 1, we also present a single analysis, a generalized mixed model (GLMM) with sites and species as random factors and including all levels of aggregation in a single analysis. This file also contains a model selection process. We used GLMM with residual maximum likelihood (REML) in the R package `lme4` (Bates and Maechler 2010; R-Core-Team 2014). As outlined above, we were not interested to study explicitly phylogenetic signals in extinction risk and did not include a complete covariance matrix on phylogenetic relations among our species in our GLMM analysis.

Independently from that, we tested whether rarity had an effect on local extinction risk using GLMM with sites and species as random effects.

Results

Effect of population size on extinction

First we analysed the effect of population size on extinction, which showed a highly significant negative effect of population size on extinction in a binomial GLM with a population size effect of -0.4869 ($SE = 0.1342$, $z = -3.630$, $p < 0.001$, $AIC = 360.84$). This result

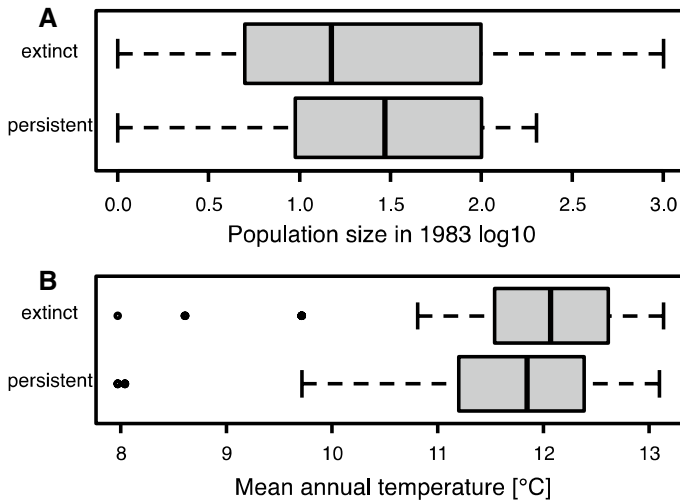


Fig. 1 Effect of population size and mean annual temperature on extinctions; note that the dependant variable extinct/persistent was used for grouping and the logarithmic scale for population size

was confirmed in a GLMM with site and species as random factor and population size as fixed factor with a slope of -0.8954 ($SE=0.2305$, $z=-3.885$, $p<0.001$, $AIC=329.8$). The larger population size the lower was the extinction risk (Fig. 1A).

Effect of mean annual temperature (MAT) on extinction

We then analysed the effect of MAT separately on extinctions in a binomial GLM, which showed a positive effect of temperature on extinction (slope 0.4354 , $SE=0.1467$, $z=2.968$, $p=0.003$, $AIC=365.6$), i.e. more extinctions in warmer sites. We also run a GLMM, including sites and species as random factor, which showed a positive effect of temperature on extinction (slope $=0.5531$, $SE=0.2315$, $z=2.389$, $p=0.017$, $AIC=344.3$). Overall, high mean annual temperature increased local extinction, with more extinctions being in warmer lowland sites (Fig. 1B).

Effect of traits and Ellenberg values on extinctions

The principal component analysis (PCA) combining all plant traits and Ellenberg indicator values showed that a number of traits and Ellenberg values were correlated across our data-set of 24 species (Fig. 2). The first three axes with a cumulative proportion of variance of 63% identified a set of plant traits explaining variance between species with the first principal component (PC1) being negatively related to plant size (PS, loading -0.51 on PC1) and seed mass (SM, -0.34) and positively to Ellenberg's temperature (T, 0.46) and moisture (F, 0.33) values, the second component being positively related to seed number (SN, loading 0.54 on PC2) and negatively to seed mass (SM, -0.44) and the third to seed bank survival (SB, loading -0.55 on PC3) and negatively to Ellenberg's nutrient value (N, -0.44).

Based on the PCA we retained the three plant traits that were most related to the first three PCA axes as uncorrelated explanatory variables, which we used in a GLM with a

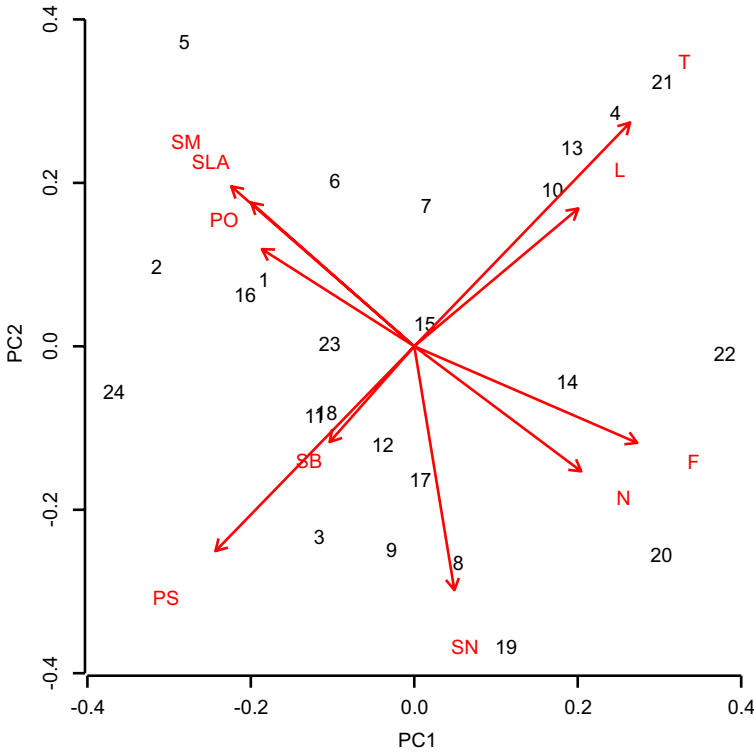


Fig. 2 Principal Component Analysis of plant traits and Ellenberg indicator values of the studied species (*SB* seed bank survival; *SN* seed number, *SM* seed mass, *PS* plant size, *SLA* specific leaf area, Ellenberg’s moisture (*F*), temperature (*T*), light (*L*) and nutrient (*N*) value, *PO* pollen:ovule ratio)

Table 2 Effects of plant size, seed number and seed bank survival on extinction in a logistic GLM; variables where standardised prior to analysis

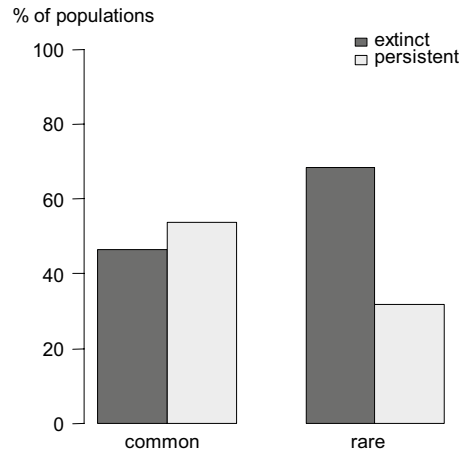
Explanatory variables	Estimate ± SE	z value	p
Plant size (PS)	0.2853 ± 0.1442	1.978	0.0479*
Seed number (SN)	− 0.3560 ± 0.1460	− 2.438	0.0148*
Survival in soil seed bank (SB)	− 0.2800 ± 0.1420	− 1.971	0.0487*

The model was extinctions/persistence ~ PS + SN + SB, where *PS* plant size, *SN* seed number, *SB* seed bank survival

logit link (logistic regression). We performed this GLM on data aggregated at the species level in order to analyse the persistence/extinction as success/failure presented in Table 2.

This GLM showed a negative effect of seed number on the persistence/extinction ratio with highly productive species being less exposed to extinctions compared to species with reduced seed production (Table 2). There were positive effects of plant size on the persistence/extinctions ratio, large plants having a higher probability of extinction than small plants (Table 2). The negative effect of survival in the soil seed bank

Fig. 3 Local extinction risk of populations ($n=290$) for 24 species according to their frequency at the regional level (GLMM, Wald's $z=3.358$, $p<0.001$)



on persistence reflects the fact that species with seed that survived well in our 2.5 year burial experiment had a lower probability of extinction of their populations (Table 2).

The GLMM analysis presented in Fig. 3 revealed that there was a highly significant effect of species' rarity status on extinction events with sites and species as random effects. Species that were rare at a larger scale showed higher proportion of local extinctions than species that were common.

Discussion

Our approach and our data highlighted for the first time that local plant population extinction is related to plant traits, niches and site conditions; with significant effects of population size, position in a climatic gradient, habitat requirements and plant traits.

One of the strongest effects on local population extinction was that smaller populations went extinct more easily than larger ones. This is additional evidence for the higher extinction risk of small populations—a well-known phenomenon but with little empirical evidence- (but see e.g. Fischer and Stöcklin 1997; Matthies et al. 2004). The higher extinction risk of smaller populations has potentially three causes: (i) higher influence of stochastic events in reproduction, growth and mortality, (ii) high impact of stochastic deterioration of the environment (Menges 1998) (iii) consequences of reduced fitness due to limited genetic diversity, caused by inbreeding, accumulation of deleterious mutations and genetic drift (Ellstrand and Elam 1993).

Additionally, there were more local extinctions of annual plants in warmer sites as indicated by mean annual temperature (MAT) in our analysis. MAT is higher in lowland areas, where summer drought is stronger and vegetation period shorter. However, lowland sites tend also to be subjected to heavier intensification of agriculture, fragmentation and urbanisation since they are close to growing urban areas. Would temperature have increased strongly, species should have disappeared in the warmest sites according to their temperature requirements. These analyses including indicator values (Ellenberg et al. 1992) suggest that changes affect lowland sites which harbour temperature demanding species, but our data did not specifically highlight temperature demanding species to disappear first.

The most important variance in plant traits explaining population extinction was related to seed number and seed mass. Populations of species with a high seed production showed decreased probability of extinction compared to plants with low seed production. Seed production was associated with the second principal component in our PCA (Fig. 2), reflecting the well known seed number—size trade-off (Moles and Westoby 2006), and negatively related to seed mass. It has repeatedly been shown that higher seed production is associated with frequent disturbances (Pakeman et al. 2008) and that large seeded plants are particularly endangered by the disappearance of effective dispersers (Poschlod et al. 2013). Our data suggest that changes in the disturbance regime and habitat of mediterranean cereal fields favour small seeded highly productive plants and that species with large seeds and low seed production are at higher risk of extinction.

Plant size was positively associated with extinction probability on our analysis (Table 2) indicating that large plants are at higher risk of extinction. This can be related to competition between arable weeds and wheat (Dutoit et al. 2001) and can be linked to the increasing stem density per square meter of cultivated cereals in our area. This suggests to use a management for less dense stands of extensive cereal fields, a causal relationship and recommendation that has already been elucidated for central Europe (Schneider et al. 1994). Density increased due to the application of manure, chemicals and herbicides that all increase productivity in cereal stands, which disadvantage plants adapted to low nutrient levels and lower cereal density (Storkey et al. 2012).

Survival of seeds in the soil seed bank was negatively associated with extinctions, with plants that had high survival in our 2.5 year seed burial experiment having lower probabilities of extinctions. This is in line with a number of theoretical (Cohen 1966; Venable and Brown 1988) and empirical works (Stöcklin and Fischer 1999; Venable 2007) showing that soil seed banks play an important role to increase local persistence and decrease probability of extinction. In our data-set, species with low seed survival in the seed bank were at highest risk of extinction in mediterranean cereal fields. Together with data on seed production and seed size, this highlights the need of special attention to short-lived and large seeded plant species such as *Agrostemma githago*, *Asperula arvensis* or *Vaccaria hispanica*, being particularly endangered after current changes in land-use.

Altogether, our data support empirically the importance of interspecific differences in plant traits for extinction risk assessment (Menges 2000; Poschlod et al. 2000) and traces a way how environmental changes and traits can be used to quantify these extinctions and predict future extinctions. We therefore propose to integrate traits e.g. in population viability analyses for plants (Brigham and Schwartz 2003; Harrison and Ray 2002; Menges 2000). However, there are a number of complex interactions between traits, population size and site characteristics which cannot be studied with such a limited set of species in a multi-population survey and which would profit from analyses using larger data-sets on extinctions.

Furthermore, there are more extinction events in rare compared to widespread species. This can be related to dispersal limitation via seeds or pollen and therefore shows the importance of nearby other populations in a metapopulation for local extinctions of plants (Harrison and Ray 2002). Only recently it was shown that specialist arable plants declined stronger than non-specialist ones (Meyer et al. 2013). However, risks that already drove species to rarity in the past on a regional scale continued to act in the observation period and lead to the observed correlation of recent extinction and regional rarity. The relative rarity of small populations of plants with high soil seed mortality at the beginning of surveys indicates that past risks influenced the relative frequencies of species at the outset. Additionally, this extends previous analyses (Fréville et al. 2007; Lavergne et al. 2006) by

focusing in a more detailed way on population extinctions and that help to identify underlying processes in order to better inform conservation efforts.

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