


Spring water deficit and soil conditions matter more than seed origin and summer drought for the establishment of temperate conifers

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Abstract In anticipation of more severe summer droughts, forestry in temperate Europe is searching for drought-resistant ecotypes of native tree species that might maintain ecosystem services in the future. We investigated how spring precipitation and soil conditions interact with summer drought and affect the establishment of conifer seedlings from different climatic origin. Emergence, establishment and subsequent performance of seedlings originating from autochthonous, Central Alpine, continental Eastern European, and Mediterranean *Pinus sylvestris* and *Picea abies* populations were studied in the dry Alpine Rhine valley, Switzerland, at three sites with differing soil water holding capacities and in 3 years with contrasting weather conditions. In addition to this natural inter-annual variation, precipitation was manipulated within sites with throughfall reduction roofs. Seedling establishment and growth were principally affected by the spring weather in the year of emergence. In years with average to positive spring water balance, seedlings grown at the site with the highest water holding capacity had 2–5 times more aboveground biomass than seedlings grown at sites with less favourable soils. Effects of seed origin were marginal and only detectable at

the drier sites: contrary to our expectations, seedlings from the Central Alpine Rhone valley, where the climatic spring water deficit is large, outperformed those from the Mediterranean. Consequently, plantation of non-native populations from dryer origin will mitigate the effects of increased summer drought at driest sites only, while the inter-annual variability of spring precipitation will continue to enable temperate conifers to regenerate on a wide range of forest soils independent of seed origin.

Keywords Climate change · *Picea abies* · *Pinus sylvestris* · Soil moisture · Tree seedlings

Introduction

Rising temperatures and increasing frequency and duration of dry spells are expected to result in a northward shift of suitable climate for European tree species (Thuiller et al. 2006; Hanewinkel et al. 2013; Lindner et al. 2014). It has been questioned, however, whether long-lived woody species will be able to keep track with rapidly changing climate and reach the sites with favourable growing conditions before they become extinct at lower latitudes (Lischke et al. 2006; Aitken et al. 2008; Renwick and Rocca 2015). Together with recurrent accounts of drought-induced tree mortality (Bigler et al. 2006; Carnicer et al. 2011; Hereş et al. 2014), this has triggered a debate about the future management of temperate European forests. Plantation of ecotypes from the dry end of current species ranges or the introduction of more drought tolerant non-native tree species has been proposed to maintain forest productivity and other ecosystem services (Kapeller et al. 2012; Hanewinkel et al. 2013; Bussotti et al. 2015). But, an extensive transplant study in the South-Eastern United States revealed that

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seedlings of potential migrant species from warmer regions did not perform better than native tree species under drier conditions (Ibáñez et al. 2009). Ecotypic differences between populations might be offset by environmental heterogeneity, which has been widely neglected in climate manipulation experiments and modelling studies. Buffering of climate change effects seems to occur in terms of soil heterogeneity (Fridley et al. 2011; Fridley and Wright 2012; García-Palacios et al. 2012; Piper et al. 2013), fine-scale thermal variability, which exceeds the expected climate warming in both topographically complex and flat terrain (Scherrer and Körner 2011; Lenoir et al. 2013; Slavich et al. 2014), as well as with respect to the tempering effects of forest canopy (von Arx et al. 2013). This is in line with Leuzinger et al. (2011), who demonstrated that effect sizes of climate change impacts diminish rapidly once models and experiments incorporate more than two factors or target a regional rather than a local scale.

Uncertainties in the projection of future tree ranges further relate to the fact that projections of precipitation changes, potentially the main driver of future tree regeneration on xeric sites in Central Europe (Richter et al. 2012), are highly uncertain with respect to magnitude as well as spatial and temporal extent (Hawkins and Sutton 2011; Fischer et al. 2015). Drought does not affect trees uniformly over the year and effects depend on site water balance. Lévesque et al. (2014) showed, for instance, that tree-ring width of *Pinus sylvestris* and *Picea abies* is negatively affected by summer drought on mesic sites, while water deficit from previous autumn to spring seems to be decisive on xeric sites. The latter corresponds well with the timing of xylem and phloem formation of the two species at lower elevations, which peaks between April and May (Swidrak et al. 2014). Not only tree growth but also tree regeneration is tied to the spring season, when seed germination takes place. Especially in smaller seeded species with limited nutrient storage such as *P. sylvestris* and *P. abies*, to be or not to be may be decided before the beginning of the summer season (Castro 2006). Given the different timing of the rainy season between the Mediterranean (winter/spring) and Central Europe (summer), Mediterranean tree species or Mediterranean ecotypes of temperate tree species may not necessarily be better adapted to the future climatic conditions projected for Central Europe, as has been suggested based on experiments studying summer drought (Richter et al. 2012; Taeger et al. 2013; Thiel et al. 2014).

We tested whether ecotypes of *P. sylvestris* and *P. abies* from the dry end of both species' distribution ranges establish more successfully than autochthonous populations under the future climatic conditions of the dry Alpine Rhine valley, Switzerland. Germination and early seedling establishment of autochthonous and non-autochthonous populations from the Central Alpine Rhone valley (no distinct

rainy season), continental Eastern Europe (rainy season in summer), and the Mediterranean (rainy season in the winter half-year) were studied in small clear-cuts with different soil water holding capacities and in 3 years with contrasting weather conditions. Throughfall reduction roofs were used to simulate a potential decrease in water availability during the growing season, as projected for the studied region by the end of the twenty-first century (Fischer et al. 2015).

Materials and methods

Species and seed material

Pinus sylvestris L. and *Picea abies* (L.) H. Karst. are the most widespread conifers in Europe, ranging from southern Spain (*P. sylvestris*) and the Balkan peninsula to Scandinavia and Siberia (Mátyás et al. 2003; Skrøppa 2003; Fig. 1). In Central Alpine valleys, *P. sylvestris* dominates low-elevation forests between 600 and 1300 m a.s.l., whereas *P. abies* prevails at higher elevation. In Central Europe, *P. abies* has been extensively planted outside its natural range to maximise wood production, especially at low elevations, where its longer-term persistence is questioned in the advent of climatic change (Hanewinkel et al. 2010). We collected seeds of both species from autochthonous populations on dry sites in the Rhine valley (autochthonous seed source), the Rhone valley (Central Alps), Tyrol, the Vienna basin, Transylvania (continental Eastern Europe), and, in the case of *P. sylvestris*, across the Mediterranean (Table 1; Fig. 1). Seeds from 3–5 trees per population were mixed in equal parts before sowing.

Sites and experimental setup

The experiment was set up in the Rhine valley near Chur, Switzerland, and included three forest sites located on south-facing, low-elevation slopes (750–850 m a.s.l.) with calcareous parent material (Table 2). The soils were classified as brown earth or rendzina, all with medium to moderately fine textured surface horizons (clay content between 20 and 25%) and a low to medium fine-earth bulk density of 0.65–1.26 g cm⁻³ (methods followed Walthert et al. 2004; detailed description in Moser et al. 2015). Although the soils have similar textures and fine-earth bulk densities, their water holding capacity differs considerably. While site 2 is located on compact rockslide material hardly penetrable to roots (piercing with a soil corer of 1 cm diameter revealed an average soil depth of 39 cm ± 15 cm SD; *N* = 40), the soil at site 3 is gravelly with a coarse fragment fraction of 30% in the surface horizon and >70% in the parent material horizon. Consequently, the plant available water holding capacity (AWC), calculated according

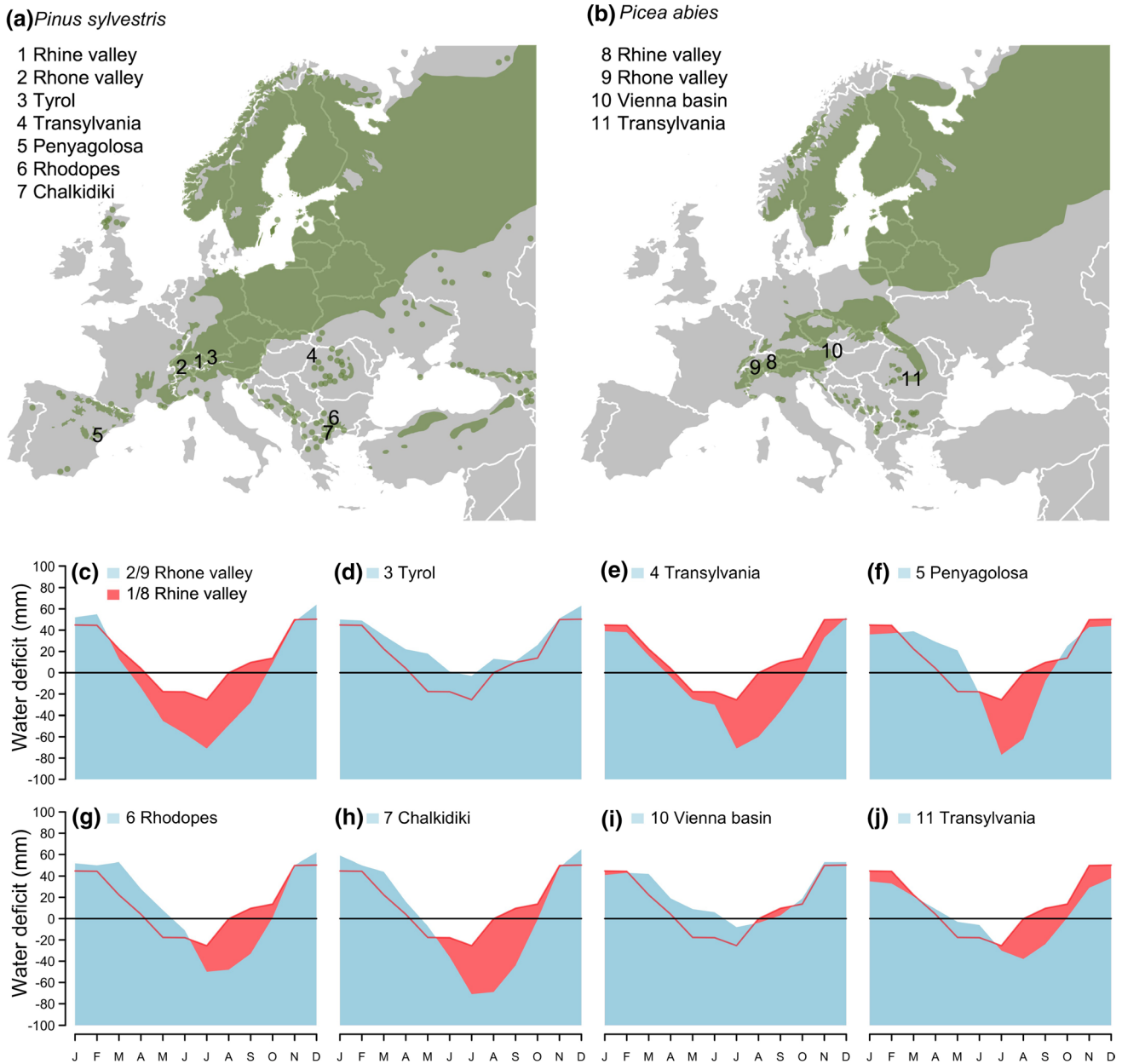


Fig. 1 Distribution of **a** *Pinus sylvestris* and **b** *Picea abies* in Europe (<http://www.euforgen.org>) and origin of the seeds used in the experiment (cf. Table 1). **c–j** Comparison of monthly climatic water deficit (mm) at the seed origin (blue) and the location of the autochthonous

population in the Rhine valley (orange). Meteorological data refer to the period 1950–2000 and were provided by MeteoSwiss (1, 2, 8, 9) and <http://www.worldclim.org> (3–7, 10, 11), respectively. A colour version of the figure is available in the online journal

to Teepe et al. (2003), was highest at site 1 and lowest at site 3 (Table 2).

The sites underwent clear cutting in an area of 0.1 ha in winter 2009/2010 to guarantee equal light conditions for all experimental units. In spring 2010, 10 sowing plots of 50 cm × 50 cm were prepared at each site by removing the vegetation and sieving the top 10 cm of the soil to 5 mm

to create a uniform seedbed. At each sowing plot, a precipitation throughfall reduction roof of 1.5 m × 1.5 m was installed, which consisted of 16 V-shaped plastic channels (Online Resource 1). The channels were mounted so that, in theory, either 100 or 33% of the precipitation should reach the soil surface. Throughfall measurements revealed that in reality 40% (±18%) of the actual precipitation

Table 1 Seed origin, climatic conditions at the seed origin (T_{Jan} average January temperature, T_{Jul} average July temperature, P average annual precipitation, WD_{AMJ} water deficit Apr–Jun, WD_{JAS} water deficit Jul–Sep), and seed quality in terms of germination rate (%) underlaboratory conditions (200 seeds incubated on filter paper at 25/13 °C day/night cycles and 90% humidity) of *Pinus sylvestris* and *Picea abies* populations included in the experiment

ID	Origin	Elevation (m a.s.l.)	T_{Jan} (°C)	T_{Jul} (°C)	P (mm)	WD_{AMJ} (mm)	WD_{JAS} (mm)	Germination lab		
								2010	2011	2013
<i>Pinus sylvestris</i>										
1	Rhine valley, Switzerland	570	0.0	17.9	810	−24	−8	97	96	94
2	Rhone valley, Switzerland	560	−0.7	19.2	603	−116	−148	96	98	97
3	Tyrol, Austria	710	−1.7	17.2	917	41	20	95	92	86
4	Transylvania, Romania	120	−2.0	20.7	609	−59	−167	–	–	42
5	Penyagolosa, Spain	1200	1.2	17.8	687	30	−147	–	–	100
6	Rhodopes, Bulgaria	1400	−3.8	15.1	664	25	−131	–	–	93
7	Chalkidiki, Greece	1360	−1.8	17.4	613	−27	−183	–	–	86
<i>Picea abies</i>										
8	Rhine valley, Switzerland	670	0.0	17.9	810	−24	−8	49	53	79
9	Rhone valley, Switzerland	620	−0.7	19.2	603	−116	−148	78	53	45
10	Vienna basin, Austria	800	−3.7	16.3	835	35	−8	61	62	63
11	Transylvania, Romania	600	−4.0	18.5	669	0	−91	–	–	75

Meteorological data refer to the period 1950–2000 and were provided by MeteoSwiss (1, 2, 8, 9) and <http://www.worldclim.org> (3–7, 10, 11)

Table 2 Location and soil characteristics of the experimental sites. Plant available water holding capacity (AWC) was calculated according to Teepe et al. (2003) and refer to 0–100 cm soil depth (site 2: 0–39 cm, according to the soil depth measurements)

	Site 1	Site 2	Site 3
Latitude/longitude	46.84/9.44	46.82/9.37	46.84/9.43
Elevation (m a.s.l.)	720	730	820
Soil type	Brown earth	Rendzina	Rendzina
AWC (mm)	145	117	75
Ammonium (mg l ^{−1})	0.44 ± 0.03	0.34 ± 0.04	0.59 ± 0.08
Nitrate (mg l ^{−1})	82.1 ± 11.4	110.2 ± 29.3	41.7 ± 8.1
Phosphate (mg l ^{−1})	<0.15	<0.15	<0.15

reached the soil surface under the 100%-roofs, and 26% (±18%) under 33%-roofs. We assume that the unexpected throughfall reduction under the 100%-roofs was due to a considerable amount of water drops sticking to the channels and evaporating before falling on the ground (interception effect). The roofs were in place from April to October each year.

The sowing plots were divided into 25 quadrats of 10 cm × 10 cm and each quadrat was sown with 15–20 seeds of one random combination of species and population in April 2010, 2011, and 2013, respectively. Some combinations of species and population were sown in 1 year only (Table 1). The sowing plots were covered by a wire cage during the first five months after sowing to prevent seed predation and herbivory from mice and birds.

Environmental measurements

Soil moisture in the topsoil was measured with matric potential sensors (MPS-1; Decagon, Pullman, USA). In 2010, measurements in four sowing plots at 15 cm soil depth showed that soil moisture varied more between sowing plots than between roof types and experimental sites. As a consequence, we continuously added sensors so that by 2014, 30 MPS-1 sensors had been installed at 15 cm soil depth and 12 sensors at 5 cm soil depth. Precipitation (ARG100 tipping bucket raingauge, Campbell Scientific, Loughborough, UK), temperature and relative humidity (EL-USB-2+ sensors, Lascar Electronics, Salisbury, UK) were measured 2 m above ground from 2010 to 2014. Several of these sensors were temporarily damaged, so that we used continuous meteorological data from a nearby meteorological station (Chur, MeteoSwiss) for data analyses. Locally measured data did not differ between sites nor from the data measured at the MeteoSwiss station (ANOVA of monthly averages April 2010–September 2014: temperature $F_{3,99} = 0.349$, $P = 0.790$; precipitation $F_{3,80} = 1.318$, $P = 0.274$). Based on these data, quarterly water deficits (precipitation – potential evapotranspiration; Thornthwaite 1948) were calculated as a measure of seasonal weather conditions (spring: April–June; summer: July–September; growing season: April–September).

Nutrient availability was estimated with ion-exchange resin bags, one of which was buried at the border of each sowing plot at 5 cm soil depth between April and November 2014. Methods followed Schleppei et al. (2012) except

that we extracted ions by shaking the resin of each bag in 80 ml 1 M KCl for 1.5 h. Ammonium was measured with flow injection analysis; nitrate and phosphate were analysed with ion chromatography after removing chloride from the solution (laboratory of the Swiss Federal Research Institute WSL, accredited according to ISO17025). Phosphate concentrations did not reach the limit of detection of 0.15 mg l^{-1} .

Seedling measurements

Seedling emergence and mortality were initially recorded bi-weekly, later monthly. Aboveground biomass (dry weight) of a random sample of 1–18 seedlings per sowing quadrat was measured at the end of the second growing season. In 2014, differences in photosynthetic activity between study sites and populations were investigated using thermal imaging (Costa et al. 2013). On a daily scale, crown temperatures of stressed and non-stressed trees are most distinctive when photosynthetic activity is reduced due to stress (Bachofen et al., unpublished data). Differences increase with the duration of the drought (Scherrer et al. 2011). In a first step, we thus analysed the diurnal pattern of assimilation by measuring stomatal conductance with leaf porometers (SC-1; Decagon, Pullman, USA) on a small number of seedlings at 1.5 h intervals between dawn and dusk. We found an assimilation peak between 9 a.m. and 1 p.m. followed by a gradual decline until dusk (Online Resource 2). Based on these data, we chose late afternoon to estimate stomatal conductance of all seedlings by thermal imaging: at each site, four thermal images were taken per plot (VarioCAM hr, InfraTec GmbH, Dresden, Germany) between 3 p.m. and 6 p.m. on a cloudless day in August 2014. Each plot contained a wet reference surface (Reinert et al. 2012) to calculate relative crown temperature $\Delta T_{\text{Wet-Crown}}$, where T_{Crown} is the minimum crown temperature of one quadrat and T_{Wet} the surface temperature of the wet reference, both determined with the IRBIS 3 software (InfraTec GmbH, Dresden, Germany; Online Resource 2). Using relative instead of absolute crown temperature allowed us to compare plots and sites even if environmental conditions differed (Scherrer et al. 2011).

Data analyses

The rate of emergence and the number of seedlings present at the end of the first growing season were standardised based on the number of sown seeds and germination rates under laboratory conditions (seed quality), i.e., emergence and number of seedlings refer to 100 viable seeds. Standardisation was necessary because seed quality varied considerably between species and populations (Table 1). Effects of site, roof type, seed origin and weather on

seedling emergence and aboveground biomass were analysed with linear mixed models (LME; R package lmerTest; R Core Team 2015). For analyses of the effect of weather on seedling establishment and seedling growth, only those populations that were sown in all 3 years were included. Effects of seed origin were studied in more detail in the 2013 cohort, which contained a larger number of populations than other cohorts and measurements also included crown temperature. Poor seed quality considerably reduced emergence of the *P. sylvestris* population from Transylvania so that this population had to be excluded from the analysis of seedling establishment. Differences in plant available water and nutrients between sites, roof types, and soil depth were analysed with analysis of variance (ANOVA).

Results

Water and nutrient availability

Weather and roof type affected soil moisture differently at the three study sites. At the site with the highest water holding capacity, average matric potential from April to September and the number of dry days (matric potential $< -100 \text{ kPa}$) at 15 cm soil depth were related neither to climatic water deficit during the growing season nor to roof type (site 1; Online Resource 3). At the other sites, however, the matric potential was lower and dry days more numerous under the 33%-roofs than under the 100%-roofs. Only at the driest site (site 3), the number of dry days was negatively related to the climatic water deficit of the growing season (Online Resource 3). The average matric potential and the number of dry days from April to September 2014 did not differ between 5 cm and 15 cm soil depth (ANOVA on 24 sensors: $F_{1,12} = 0.016$, $P = 0.900$; $F_{1,12} = 0.046$, $P = 0.834$).

Plant available nutrients measured in terms of ammonium and nitrate differed neither between study sites nor between roof types (ANOVA, $P > 0.05$; Table 2).

Weather and site effects on seedling performance

Seedling establishment involves a series of events such as seed germination, seedling emergence and survival, which result in differing seedling numbers. The number of *P. sylvestris* seedlings that had established by the end of the first growing season was first and foremost affected by climatic spring water deficit (Table 3). In years with favourable spring weather conditions, three to five times more seedlings emerged and survived than in years with a dry spring season (Fig. 2a). Establishment also depended on study site (Table 3), with a 1.5 to 3-fold higher number of seedlings at site 1 (high AWC) than site 3 (poor AWC).

Table 3 Results of linear mixed effects models for the effects of study site, roof type, seed origin as well as spring (Apr–Jun) and summer (Jul–Sep) water deficit in the in the year of seedling emergence ($\Delta\text{WD}_{\text{AMJ}}$ site 1st year and $\Delta\text{WD}_{\text{JAS}}$ site 1st year, respec-

tively) and in the second growing season ($\Delta\text{WD}_{\text{AMJ}}$ site 2nd year and $\Delta\text{WD}_{\text{JAS}}$ site 2nd year, respectively) on the standardised number of seedlings at the end of the first growing season and aboveground biomass at the end of the second growing season (details s. Fig. 2)

Source of variation	<i>Pinus sylvestris</i>						<i>Picea abies</i>					
	No of seedlings			Biomass (g) ^a			No of seedlings			Biomass (g) ^{a,b}		
	df	F	P	df	F	P	df	F	P	df	F	P
Study site	2,26	13.8	<0.001	2,24	13.6	<0.001	2,26	22.9	<0.001	–	–	–
Roof type	1,26	0.3	0.572	1,23	2.6	0.123	1,26	7.6	0.010	1,48	2.8	0.100
Seed origin	2,236	1.5	0.229	2,150	2.7	0.072	2,236	4.7	0.010	2,48	0.8	0.435
$\Delta\text{WD}_{\text{AMJ}}$ 1st year	1,236	136.6	<0.001	1,154	55.3	<0.001	1,236	16.5	<0.001	1,48	47.7	<0.001
$\Delta\text{WD}_{\text{JAS}}$ 1st year	1,236	7.4	0.007	1,157	1.5	0.226	1,236	0.3	0.561	1,48	0.0	0.954
$\Delta\text{WD}_{\text{AMJ}}$ 2nd year	–	–	–	1,162	0.0	0.917	–	–	–	1,48	0.1	0.788
$\Delta\text{WD}_{\text{JAS}}$ 2nd year	–	–	–	1,164	0.2	0.631	–	–	–	1,48	1.4	0.246

Bold indicate P values <0.05

^a Log-transformed

^b Only site 1 included in the model because low seedling emergence made the experimental design unbalanced at the other sites

A small climatic water deficit during summer increased survival but its effect on seedling establishment was considerably smaller than that of climatic spring water deficit and site characteristics, as indicated by the substantially lower F value (Table 3). At site 1, more than 20 out of 100 viable seeds successfully established in all years, while at the other sites this number was reached only in years with average to wet spring conditions. The aboveground biomass of 2-year-old seedlings was still affected by the climatic spring water deficit in the year of germination but not by later weather conditions, i.e., neither summer water deficit in the year of germination, nor spring or summer water deficit during the second growing season (Table 3). Site effects were also pronounced, though less important than spring weather (fourfold smaller F value, Table 3), and differences between sites mirrored those found with respect to seedling establishment (Fig. 2b). We found no differences in seedling numbers or aboveground biomass between roof types and populations (Rhine valley, Rhone valley, Tyrol).

In *P. abies*, seedling establishment followed similar patterns as in *P. sylvestris* only that site and spring weather effects were of similar magnitude (Table 3). Although establishment success varied between populations (Rhine valley, Rhone valley, Vienna basin), we found no consistent differences between populations with respect to sites and/or spring weather conditions (Fig. 2a). Seedling numbers were almost twice as high under the 100% roofs (30 ± 4 SE) than under the 33% roofs (16 ± 3 ; Table 3) despite the fact that soil moisture did not differ between the two roof types (Online Resource 3). Overall, *P. abies*

only reached sizeable numbers of seedlings at the site with the highest AWC (site 1, Fig. 2b). Consequently, the experimental design was unbalanced and analyses of second year biomass were possible at site 1 only. Consistent with *P. sylvestris*, the aboveground biomass depended on the spring water deficit in the year of germination but not on later weather conditions (Table 3). Although not statistically verified, differences in seedling biomass between study sites resembled those of establishment success (Fig. 2).

Populations from drier climates

The performance of autochthonous populations compared to those from the Central Alps, Eastern Europe or the Mediterranean was studied in more detail in the 2013 cohort, which included seven *P. sylvestris* and four *P. abies* populations (Table 1). The number and aboveground biomass of *P. sylvestris* seedlings were again mainly driven by site characteristics (LME, $F_{2,227} = 17.0$, $P < 0.001$; $F_{2,22} = 33.5$, $P < 0.001$) and differences between populations were small (LME, $F_{5,135} = 2.7$, $P = 0.022$; $F_{6,101} = 2.8$, $P = 0.015$). Neither establishment nor biomass was related to climatic parameters that potentially indicate local adaptation to summer drought, such as mean July temperature and precipitation or climatic water deficit during summer (Online Resource 4). But, biomass of seedlings growing under reduced precipitation (33%-roof type) at sites 2 and 3 was marginally higher in populations from locations with a negative climatic spring water balance than those from locations with a positive balance (Fig. 3a).

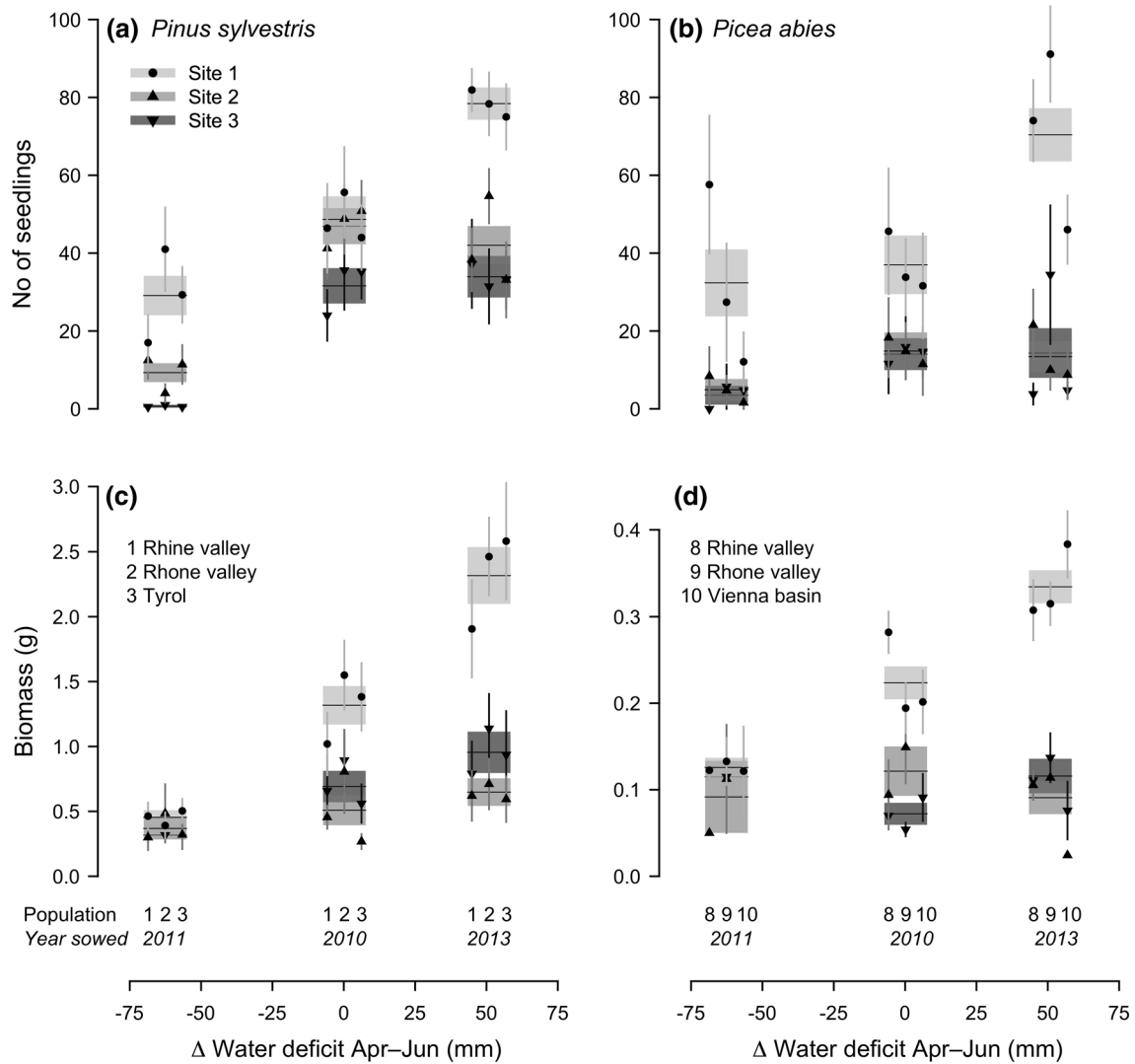


Fig. 2 Establishment and growth of *Pinus sylvestris* and *Picea abies* seedlings in relation to weather at the time of emergence, site and seed origin: **a, b** standardised number of seedlings at the end of the first growing season; **c, d** aboveground biomass at the end of the second growing season (*symbols/vertical bars* population means \pm SE;

horizontal bars/polygons site means \pm SE). Weather during the three months following seed sowing (Δ water deficit Apr–Jun) was defined as the deviation of the water deficit from the long-term average in the Rhine valley (-39 ± 61 mm SD). Only populations that were sown in all years are shown

At site 3, this corresponded to estimates of stomatal conductance that show marginally higher transpiration and concomitant lower crown temperature in seedlings from locations with a negative climatic spring water balance (Fig. 3b).

Due to the limited number of *P. abies* populations in the 2013 cohort, correlations with climatic parameters at the seed origin are rather arbitrary. Nevertheless, we found a higher number of established seedlings in the Rhone valley population, the seed origin with the most negative climatic water balance in spring, than in the population from the Vienna basin with a positive April–June water balance (LME, $F_{3,32} = 2.9$, $P = 0.049$; Fig. 4).

Discussion

Seasonal water availability and soil heterogeneity

Studying seedling establishment in several years with contrasting weather conditions demonstrated a huge impact of spring weather not only on seedling emergence but also on longer-term seedling performance. Climatic water balance during very early establishment outweighed effects of later weather conditions and seed origin (Table 3), which is not surprising since germination requires moist conditions and emerging seedlings are prone to desiccation until they have established an adequate root system (Castro 2006;

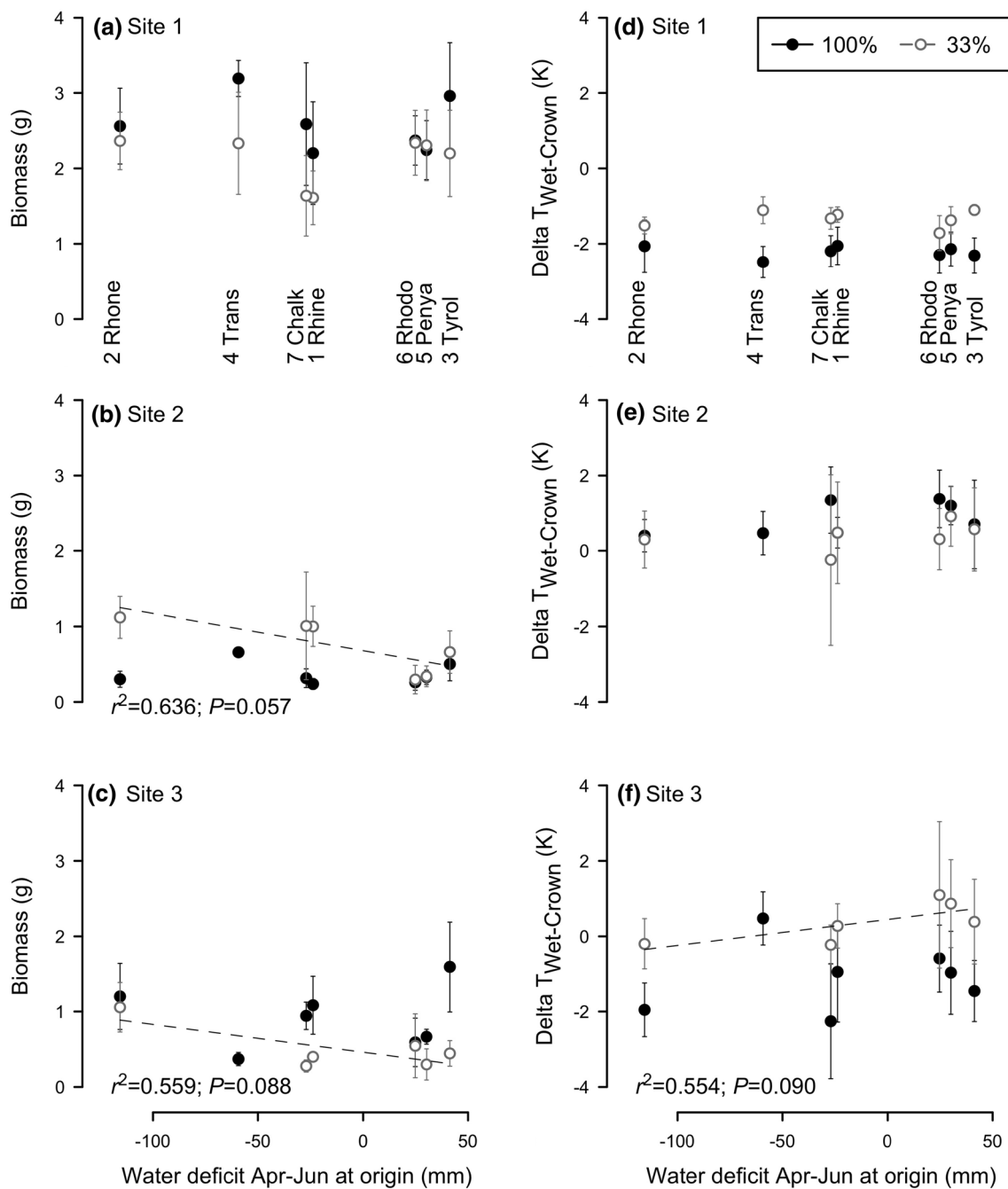


Fig. 3 Performance of 2-year-old *Pinus sylvestris* seedlings sowed in 2013 depending on study site, roof type and average spring water deficit at the seed origin (s. Table 1): **a–c** aboveground biomass (mean \pm SE); **d–f** leaf conductance measured as the thermal differ-

ence between the minimum crown temperature (T_{Crown}) and a wet artificial reference surface (T_{Wet} ; mean \pm SE). The lower $\Delta T_{Wet-Crown}$, the higher the leaf conductance of a seedling

Dobrowski et al. 2015). Environmental conditions during emergence and early establishment did not only define the number of seedlings that emerged and survived in a particular year but also affected their size (Fig. 2). The observed drought-induced reduction of seedling size was still noticeable 2 years after emergence, potentially hampering the seedlings' ability to compete with surrounding vegetation.

The number of seedlings that were able to establish in the driest year was so low and their size up to 2 years after emergence was so small that they are unlikely to contribute to long-term forest regeneration. This is in accordance with the facts that tree regeneration occurs in pulses (Grubb 1977; Zackrisson et al. 1995; Savage et al. 1996) and juveniles occupy a moister niche than adults in water limited

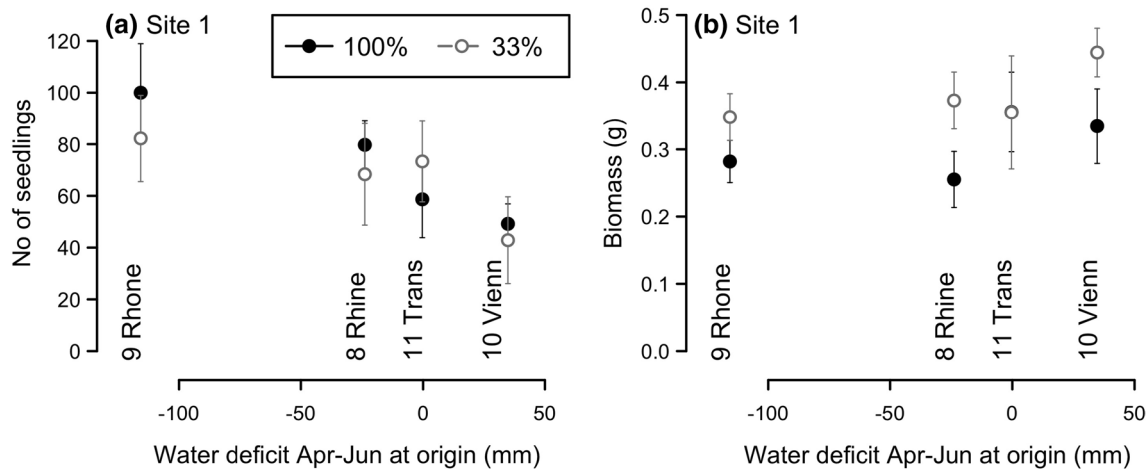


Fig. 4 **a** Number of *Picea abies* seedlings at the end of the first growing season in 2013 and **b** their average biomass at the end of the second growing season (\pm SE) at study site 1 in relation to average climatic spring water deficit at the seed origin (s. Table 1)

forest systems (Dobrowski et al. 2015). In *P. sylvestris*, the effect of climatic spring water balance on seedling abundance and size was amplified by soil characteristics with a fivefold difference in seedling biomass between the driest and the wettest year at site 1 (high AWC) and a threefold difference at site 3 (low AWC; Fig. 2). Since precipitation, temperature and nutrient availability were similar at all sites, we attribute these site effects to AWC, the sole characteristic that differentiates them. Low AWC indicates high percolation of precipitation and thus rapid drying out of the soil, whereas high amounts of fine earth at site 1 might facilitate the capillary rise of soil water from the subsoil to the topsoil. The latter might also explain why the throughfall reduction roofs had no effect on soil moisture at site 1. It remains unclear, though, if larger differences in actual precipitation reduction between the two roof types would have resulted in similar soil moisture conditions at the drier sites. The high proportion of rainfall intercepted by the 100%-roofs is unfortunate but was compensated by the high inter-annual variation in natural precipitation, which by chance included an average, a dry and a wet year, corresponding roughly to the mean, the mean negative SD, and the mean positive SD, respectively, of the long-term spring precipitation in the Rhine valley (Online Resource 5). Based on this inter-annual variation in precipitation, we conclude that both species are able to regenerate in all but dry years on soils with high AWC. By contrast, successful seedling establishment on low AWC soils is tied to exceptionally wet spring weather. The latter particularly holds for *P. abies* and mirrors the species' ecological niche, which does not extend to environments as dry as those colonised by *P. sylvestris* (Ellenberg 1988). Although climate models project a decline of summer precipitation in Central Europe by up to -40% by 2100 (CH2011 2011), our

results suggest that huge inter-annual variability of weather conditions will continue to be the basis of long-term forest regeneration (Grubb 1977) in large parts of both species' natural distribution ranges, and only a small proportion of conifer forests might become vulnerable to climate change driven regeneration failure. This is corroborated by projections of spring precipitation, which seems to be less likely to change in the future than summer precipitation (Hawkins and Sutton 2011). A comparison of climate data shows that already today the spring water balance is lower in the Central Alpine Rhone, Aosta and Venosta valleys than in the Rhine valley, where our experimental sites were located (Online Resource 5). Even in the event of a 3 K temperature increase during the spring season, as, e.g., projected for lowland Switzerland by the end of the twenty-first century based on the A2 and A1B climate change scenarios (CH2011 2011), the spring climate of the Rhine valley will, on average, remain moister than today's climate in the Rhone and Aosta valleys, where *P. sylvestris* and *P. abies* have both persisted to date. In the Rhone valley, increased mortality of *P. sylvestris* has been observed at low elevations after periods of consecutive dry years (Rigling et al. 2013). Yet, this species is known for its resilience after drought (Eilmann et al. 2010) and it has been suggested that currently observed changes in *P. sylvestris* stands are the result of an intricate mixture of drought stress, stand age, stand density (Giuggiola et al. 2013), and land-use changes (Gimmi et al. 2010).

Populations from drier climates

The uniform projections of rising temperatures and declining summer precipitation fostered the idea that drought adapted ecotypes of temperate tree species might be better

suiting to the future climatic conditions in temperate Europe than autochthonous populations (Kapeller et al. 2012; Hanewinkel et al. 2013; Bussotti et al. 2015). Local adaptation to summer drought has been found in all major European tree species (*Fagus sylvatica*: Sánchez-Gómez et al. 2013; Thiel et al. 2014; *Picea abies*: Modrzyński and Eriksson 2002; *Pinus sylvestris*: Richter et al. 2012; Taeger et al. 2013). Nevertheless, ecotypic differences between populations in terms of photosynthetic activity and seedling biomass were small compared to the effects of spring weather and site characteristics, and they were restricted to very dry conditions (site 3 with low AWC \times 33% roof; Fig. 3). Under these circumstances, the climatic water deficit at the origin during spring rather than during the summer season was related to seedling performance. The suitability of different populations to future climatic conditions will thus depend as much on the seasonal timing as on the intensity of future droughts. Our study suggests that Central Alpine populations of temperate European conifers are likely to be better adapted to the future climatic conditions in Central Europe than Mediterranean populations, because they are adapted to highly variable spring weather and a lower climatic spring water balance (Fig. 1). In the Mediterranean, by contrast, water is generally abundant during winter until late spring, which enables seeds to germinate and establish before the onset of the summer drought (Castro et al. 2004; Castro 2006).

Conclusions

Regeneration of *P. sylvestris* and *P. abies* in dry Alpine regions such as the Rhine valley is tied to years with favourable spring weather conditions. During such windows of opportunity, the majority of seedlings are able to establish, independent of their origin. Even under changing climatic conditions, autochthonous populations will thus regenerate as successfully as non-native populations from more drought prone regions given that the frequency of years with favourable weather conditions does not decline dramatically. Tree regeneration pulses might become less frequent, because on the one hand rising temperatures will increase the climatic water deficit even if spring precipitation should remain constant (Hawkins and Sutton 2011), and on the other hand tree recruitment also depends on seed availability, which varies tremendously between years (Nussbaumer et al. 2016). The concurrence of mass seed production (masting) and favourable weather conditions might be particularly important after large-scale disturbance events such as windthrow or fire (Moser et al. 2010). Nonetheless, our study supports the perception that landscape heterogeneity is likely to buffer the effects of changing climatic conditions in many ecosystems in some

ways and to some extent (Fridley et al. 2011; Scherrer and Körner 2011; Fridley and Wright 2012; García-Palacios et al. 2012; Lenoir et al. 2013; Piper et al. 2013; von Arx et al. 2013; Slavich et al. 2014). It suggests that considerable buffering potential not only exists with respect to soil heterogeneity and topography, but also regarding the timing of regeneration and growth processes, which might (Laube et al. 2014), but not necessarily, be affected by climatic change.

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Author contribution statement TW and BM conceived and designed the experiment. BM, TW, MM, UW, and LW conducted the fieldwork. BM analysed the data, LW collaborated in soil analysis. BM wrote the manuscript with editorial advice from other authors.

Compliance with ethical standards

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Conflict of interest The authors declare that they have no conflict of interest.

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