Towards an understanding of factors controlling seed bank composition and longevity in the alpine environment

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I. Abstract

The ability of seeds to regenerate from soil seed banks has long been recognized as a key survival strategy for plants establishing new niches in highly variable climates of alpine environments. However, the fundamental aspects of evolutionary/selective forces for seed bank development in alpine ecosystems are largely unknown. Here, we developed a model that describes dormancy, a high temperature requirement and a specific light/darkness regime at the time of seed shedding can preclude autumn germination, thus contributing to seed persistence until the next growing season. The benefits of these factors synchronising germination with the growing season are reviewed. Additionally, the importance of climatic variations of maternal environment affecting some of these factors is also discussed. It is suggested that the environmental conditions during the growing season partly control the seed persistence and seeds that fail to germinate are carried over to the next season. Species that have small (<3 mg) and round-shaped seeds tend to persist more easily in soil for over five years, than do the large or flat seeds. However, some large-seeded species also have the potential to establish short-term persistence bank. A literature survey reveals 88% of the alpine seeds have a mass <3 mg. Seed size has only a weak relationship with mean germination timing (MGT) indicating that reduced persistence in large-seeded species cannot be counteracted by quicker germination, but combined effects of other factors stimulating germination remain an open area to be studied. It is proposed that long distance dispersal (LDD) is limited in most-but not all-species, primarily due to the absence of specialized dispersal structures. However, among numerous dispersal modes, most species tend to be dispersed by wind. Thus, spermatophytes of alpine environments have a greater tendency to establish seed banks and spread the risk of germination to many years, rather than being dispersed to other micro-climates.

Key words: Alpine tundra, climate change, dormancy, germination, phylogeny, seed size, seedling establishment, light requirement, seed dispersal.

II. Introduction

The ecological significance of seeds in establishing plant communities has been extensively documented for an appreciable number of alpine species (Bliss, 1958; Baskin and Baskin, 1998; Molau and Larsson, 2000; Welling and Laine, 2000; Welling *et al.*, 2004; Bu *et al.*, 2009; Ma *et al.*, 2011). Unlike other ecosystems, alpine seed germination is principally determined by the short and erratic growing season (Bell and Bliss, 1980). Consequently, seeds do not often have a favorable germination environment and would be expected to enter the soil seed bank to postpone the germination to a later year when more favorable conditions occur (reviewed by Arroyo *et al.*, 1999; Cavieres and Arroyo, 2001; Welling *et al.*, 2004). However, there is some evidence to show that the unfavorable conditions preventing seed germination may sometimes persist for two consecutive years (Körner, 2003). Therefore, it is not surprising to discover that evolutionary and ecological processes have resulted in most, if not all, species adapted to alpine regions having seeds with a longevity of at least two years.

The ability of alpine seeds to form long-term seed banks is considered as an important ecological trait for population dynamics and species establishment (McGraw and Vavrek, 1989; Körner, 2003). One of the important means by which seeds persist in the soil is by preventing germination immediately after dehiscence. Early efforts to understand alpine soil seed bank dynamics have mostly focused on the seed germination timing *per se* (see Baskin and Baskin, 1998), thus the mechanisms controlling germination have not been treated adequately. Even less attention has been paid to recognize the factors influencing seed persistence in alpine environments. This dearth of knowledge has become an acute problem, especially given that the effects of global warming conspicuously affect the alpine ecosystem as a whole rather than impacting upon specific species (Walther *et al.*, 2002; Gottfried *et al.*, 2012). Such predictions explicitly require a profound understanding of seed adaptation and it is imperative to model the imminent alpine flora community changes that may result from climate shifts.

The important goal of this review is to explore some of the mechanisms evolved in alpine species that determine how seeds enter and leave the soil. Our model depicts dormancy, a requirement for relatively high temperatures and specific light/darkness regimes are the key factors affecting seed persistence in alpine environment. Moreover, when considering seed persistence in an ecological context, it is imperative to place special emphasis on the comparative biology of alpine plants including the phenology of seed size, germination traits and dispersal affecting the seed turnover and contributing to persistence. We do not mean to imply that only these factors are the pertinent adaptive traits for seed persistence in alpine species; however, our view is that many unknowns impede our understanding and this oversight can be only resolved by paying more attention to these areas. We aim to identify these critical gaps and suggest possible future research directions, rather than summarizing the overwhelming information available on soil seed banks per se, a topic covered in myriad literature (e.g. McGraw and Vavrek, 1989; Körner, 2003; Baskin and Baskin, 1998). To this end, the focus of discussion is limited only to seed bank establishment and persistence. Thus, we do not address topics such as seed density in relation to seed numbers, and species competition during emergence etc. Nevertheless, understanding these subjects is vital, yet out knowledge is rudimentary, but somewhat detailed information can be

found elsewhere (see Chambers *et al.*, 1991; Chambers and MacMahon, 1994; Forbis, 2003; Körner, 2003; Bu *et al.*, 2007a; Bu *et al.*, 2008; Bu *et al.*, 2009).

(1) Definitions and terms

At the outset, it is necessary to explain our terms. Here we consider a seed to be the product of sexual reproduction dispersed at natural dispersal time with all its dispersal structures, e.g. wings, plumes etc. We do not use specific terms such as achenes, and refer to all these propagules as seeds. We also adopt the seed bank classification scheme of Bakker et al. (1996) and Thompson et al. (1997) on a broader level. Hence, seeds that persist in soil for less than 1 year are termed as 'transient'. Whereas, seeds that persist for more than 1 year but less than 5 years are characteristically considered as 'short-term persistent' and any species that continue to persist in soil for more than 5 years are 'long-term persistent'. Although, in most of the discussion, the term 'transient' and 'persistent' are loosely used to differentiate the seed longevity of less than or more than 1 year respectively, whenever possible, clear distinctions of seed bank types are made. We define dormancy as a state in which seed germination does not take place even when the favorable environmental factors (temperature, moisture, light/dark etc.) that stimulated germination in a non-dormant or dormancy broken seed are provided. Five types of dormancy have been recognized in seeds (Baskin and Baskin, 1998; Baskin and Baskin, 2004) namely physiological (PD), morphological (MD), morpho-physiological (MPD), physical (PY) and combinational (PD+PY), we attempt to specify dormancy types when information is available. For the purpose of this review, we define 'long-distance dispersal' (LDD) as seed movement which is over 100 m (sensu Cain et al., 2000).

I. Dormancy

Dormancy is one of the remarkable traits evolved in seeds that synchronize germination with the most favorable season for seedling survival (Vleeshouwers *et al.*, 1995; Baskin and Baskin, 1998; Thompson, 2000; Bewley *et al.*, 2013). Since Amen (1966) reviewed the extent and role of seed dormancy in alpine species, there has been an abrupt increase in the number of studies documenting alpine dormancy from species to community level (Baskin and Baskin, 1998; Liu *et al.*, 2011). This increase in literature has refuted two of the early notions that (i) dormancy is a rare phenomenon in alpine systems (see Amen, 1966) and (ii) dormancy is highly associated with seed-coat inhibition (Billings and Mooney, 1968). The literature available on alpine seed dormancy is in fact growing and this topic has been comprehensively reviewed periodically (Billings and Mooney, 1968; Kaye, 1997; Baskin and Baskin, 1998). It is beyond the scope here to present such an exhaustive review yet again. Instead, we discuss the role of dormancy (and breaking requirements) in seed persistence.

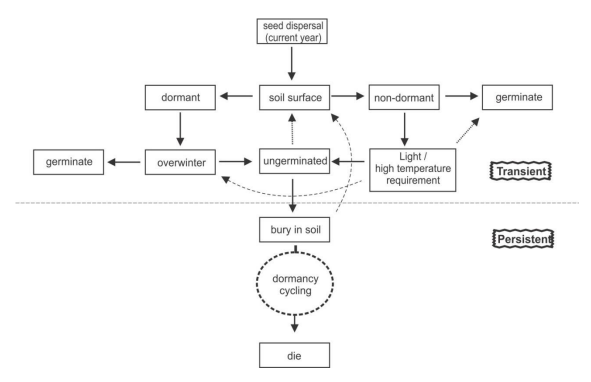


Fig. 1 Fate of a seed after dispersal from the mother plant. The horizontal dotted line distinguishes the transient and persistent bank. If the seeds buried in soil (or close to the surface) germinate or die within five years, they are transient seed banks. The remainder develop persistent seed banks. Seeds can die or be dispersed to other locations at any stage of this model (see also Chambers and MacMahon, 1994). Similarly, the soil surface may have increased seed rain, due to dispersed seeds from other locations reaching the local micro-site. For the purpose of this model, 'soil surface' means the buried depth at which seeds can receive dormancy breaking cues and/or light, germinate and establish seedlings. Seeds buried in the soil undergo seasonal cycling in dormancy status and the depth of the dormancy changes with season.

Cavieres (1999) and Cavieres and Arroyo (2001) postulated that dormant seeds have some advantage in establishing persistent seed banks in temporally and spatially unpredictable alpine environments. This is partly because seeds shed in the autumn are mostly dormant and require a period of cold-stratification satisfied under snow before they germinate in spring (see below) and partly due to the ability of the seeds to re-enter dormancy (in the case of PD) if they did not germinate. This view is in fact supported by (i) Rees' (1996) assertions that in order to maximize the progeny number, species adapted to an environment that varies from one year to another select dormancy as an adaptive trait (also see Rees and Long, 1992) and (ii) as Bewley *et al.* (2013) recently generalized, the number of species with dormant seeds is likely to increase with poleward movement because the unpredictability in climate increases.

Apart from these simple theoretical considerations, direct evidence delineating whether or not dormancy contributes to seed persistence in alpine regions is scarce. This is ascribable to the paucity of information amassed on a specific species for which both the dormancy and longevity are known. Some progress has been made recently by Schwienbacher *et al.* (2011) to bridge this gap (but see conclusion section). In an attempt to investigate the seed dormancy types in 28

alpine species of central European Alps, Austria, those authors concluded 26 species were dormant and only 2 species were non-dormant. Out of 26 dormant species, PD (with all subclasses namely non-deep, intermediate and deep) was present in 20 species. MD and PY were recognized in 4 and 2 species respectively. In addition to assigning the type of dormancy, their study included 9 species for which the seed longevity had previously been assessed by artificial burial method (Schwienbacher *et al.*, 2010). The findings of their study clearly indicate that all seeds survived for at least two years in an artificial burial experiment (see 'Seed size and persistence', below) and had some type of dormancy - mostly PD or PY (Table 1).

The assertion that dormant seeds persist in soil is in marked contrast with Thompson *et al.* (2003) who concluded that "dormancy is neither a necessary nor a sufficient condition for the accumulation of a persistent seed bank, and there is no realistic prospect of predicting persistence from dormancy" based on their data from north-west Europe (Thompson *et al.*, 1997). They further supported this conclusion by showing that most of the temperate shrubs have deep dormancy, but these species rarely persist in soil. However, we presume that two different issues have been raised and these can be encapsulated in the questions (1) do seeds of species establishing persistent seed banks have dormancy and (2) do all dormant seeds enter persistent seed banks? Thus, not all dormant seeds persist in alpine soil, but being non-dormant could reduce the ability of seeds to enter persistent seed bank in alpine environments. For example, despite *Geum reptans* and *Oxyria digyna* being dormant (Table 1), they were both short-term persistent corroborating the claim that some dormant seeds are also short-lived in soil.

Indeed, it was initially thought that factors such as light requirements, germination inhibitors *etc.* cause dormancy in alpine seeds (Amen, 1965). Consequently, Billings and Mooney (1968) concluded 'seed dormancy in most alpine species is under environmental control'. In the central European Alps, however, Schwienbacher *et al.* (2011) explicitly classified the dormancy type in accordance with Baskin and Baskin (2004), therefore these species do not represent 'enforced dormancy'. One plausible reason for the disagreement observed between North West Europe and central European Alps datasets may lie in the geographical distribution of the species. In the alpine environment, many species (ca. >70% as estimated by Baskin and Baskin, 1998 cited in Schwienbacher *et al.*, 2011) produce dormant seeds. Further evidence to safely conclude that most alpine seeds are likely to produce dormant seeds comes from the data set assembled by Jurado and Flores (2005) which suggests plants growing in environments with frost and/or drought contain some form of seed dormancy (PD, MPD and PY) than species in more benign environment.

It is generally agreed that seeds dispersed in autumn are held in seed banks until spring because of dormancy breaking requirements (Fig. 1, Table 2). These dormant seeds overwinter under snow cover and alleviate dormancy mostly by cold-wet stratification. They then germinate at the first opportunity immediately after the snow melt in spring to maximize growing periods in favorable conditions (Baskin and Baskin, 1998). Germination of non-dormant seeds in autumn, on the other hand, is exclusively controlled by the local microclimate of a particular year. Interannual variation in autumn temperature should therefore impose selective pressures on alpine communities conferring advantages on species with dormant seeds (Rees and Long, 1992; Allen and Meyer, 1998).

The influence of environmental conditions acting as selective force in the evolution of seed dormancy is gaining recognition. Studies focusing on alpine species, e.g. *Chenopodium bonushenricus* (Dorne, 1981), *Artemisia tridentata* ssp. *vaseyana* (Meyer and Monsen, 1991), genus *Penstemon* (Meyer and Kitchen, 1994b; Meyer *et al.*, 1995), *Linum perenne* (Meyer and Kitchen, 1994a) *Phacelia secunda* (Cavieres and Arroyo, 2001), and *Koenigia islandica* (Wagner and Simons, 2008), revealed seeds collected from lower altitudes are non-dormant or conditionally dormant, seeds from mid-altitudes are dormant and require cold-stratification for dormancy alleviation, and seeds from high-altitudes are dormant and require longer cold-wet stratification duration compared to seeds collected from mid-altitudes. Insights gained from these studies suggest dormancy and longer stratification duration must have co-evolved in seeds adapted to higher altitudes. The differential effects of environmental conditions on dormancy may also explain why *Oxyria digyna* has been classified both as dormant (Stöcklin and Bäumler, 1996; Schwienbacher *et al.*, 2011) and non-dormant (Mooney and Billings, 1961; Mondoni *et al.*, 2012). This species is common to both arctic and alpine communities and therefore is subject to a range of conditions which may or may not induce dormancy.

The opposite situation was observed in some of the recent studies which showed seeds of many alpine plants lack dormancy when they develop and therefore germinate immediately in autumn. Seed germination experiments performed in 37 endemic rare or endangered taxa of Sierra Nevada high mountain (SE Spain), for instance, showed most of the species tested were nondormant and do not require any pretreatment for germination (Lorite et al., 2007). Giménez-Benavides et al. (2005) reached a similar conclusion when working with 20 species collected from high altitudes of Spain. It is possible that these areas experience Mediterranean climate and seeds develop in a slightly warmer temperature than in true alpine locations, to be less dormant (see also conclusion). One recent study by Sommerville et al. (2013) shows species with nonendospermic seeds did not require any stratification to germinate, however, endospermic seeds needed 8 weeks of cold stratification for germination. Although they concluded that 'alpine species with endospermic seed and a restricted distribution are most likely to contract in range under climate change and would be appropriate to prioritize for ex-situ conservation', nonendospermic seeds are also at risk in warming climates as the absence of dormancy/breaking requirement impose the risk of autumn germination and seedling mortality in winter, particularly on exposed sites (see Mondoni et al. 2012).

Dormancy has generally not been specifically addressed in studies of seed persistence in the soil. Nevertheless, the available evidence makes it tempting to conclude that dormancy plays a crucial role in persistence at least for alpine vegetation. However, it is too early to generalize at ecosystem level. This is mainly due to the species-specific breaking requirements evolved in spermatophytes adapted to alpine environment (Table 2). Given most species with PD, these seeds can cycle back to dormancy if the conditions are unfavorable during early spring. Such cycling is also expected to be common in buried seeds (Baskin and Baskin, 1998). The evolution of a dormancy cycle, regardless of the seed position in soil, can be viewed as a specific control in preventing germination outside growing season. For example, if the buried seeds are brought back to the soil surface (Fig. 1), dormancy cycling checks the germination time to synchronize with growing season. Supposing such seasonal change of dormancy had not evolved in alpine species, there would be a chance for buried seeds (when they reach soil surface) to germinate any time of the year, provided other environmental conditions are satisfied.

Table 1 List of species for which soil persistence and dormancy type have been estimated [Adapted from (Schwienbacher *et al.*, 2010; Schwienbacher *et al.*, 2011)]. Nomenclature used in the original works was as such retained. For the abbreviations of dormancy types see text. Seed size was calculated using digitizer option in origin 8.6.

Species	Family	Seed mass (mg)	Seed bank type	Dormancy type
Achillea moschata	Asteraceae	0.21	Short-term Persistence	PD
Artemisia genipi	Asteraceae	0.25	Short-term Persistence	PD
Anthyllis alpicola*	Fabaceae	2.41	Long-term Persistence	PY
Geum reptans	Rosaceae	0.54	Transient	PD
Linaria alpina	Veronicaceae	0.14	Long-term Persistence	PD
Oxyria digyna	Polygonaceae	0.55	Transient	PD
Saxifraga aizoides*	Saxifragaceae	0.04	Long-term Persistence	PD
Saxifraga oppositifolia*	Saxifragaceae	0.08	Short-term Persistence	PD
Trifolium pallescens*	Fabaceae	0.60	Long-term Persistence	PY

^{*}Classified as transient by Cerabolini et al. (2003)

Table 2. Species that exhibit dormancy at the time of dispersal in autumn and the conditions shown to alleviate initiate dormancy and progress seeds towards germination. Seeds form 'transient' seed bank break dormancy by cold-wet stratification/after-ripening conditions in the soil. The ability of these non-dormant seeds to germinate the following spring is largely determined by environmental conditions. Data on the seed longevity of these species to differentiate short-term or long-term persistent bank are currently unavailable.

Species	Key results	References
Carex pendula L. and C.remota L.	Seeds stratification at low temperature alleviates dormancy effectively than higher temperature. Seeds of <i>C. pendula</i> and <i>C. remota</i> did not germinate below 9 and 15°C respectively. Germination requirements and dormancy patterns ceased germination until late spring and summer.	Brändel and Schütz (2005)
Fritillaria tubiformis subsp moggridgei	Seeds are dormant (MD) when dispersed in autumn, but low temperature (4°C) experienced in winter breaks dormancy and seeds readily germinate in spring.	Carasso <i>et al.</i> (2012)
Aquilegia nivalis, Lagotis cashmeriana and Meconopsis latifolia	Seeds of these three perennial species require longer period of pre-chilling during winter to break dormancy and germinate in following spring. Pre-chilling coupled with GA_3 application stimulated germination in all three species, thus supporting overwintering is mandatory for seed germination and germination is precluded in the autumn.	Dar et al. (2009)
Centaurium somedanum	Non-deep MPD in seeds controls the germination timing in alpine climate. Seeds come out of dormancy early in spring readily germinate at 15-22°C, allowing the successful seedling establishment in late spring and summer.	Fernandez-Pascual <i>et al.</i> (2012)
Sempervivum vicentei subsp. paui, Ranunculus ollissiponensis subsp. alpinus, Veronica fruticans subsp. cantabrica and Luzula hispanica.	Germination in freshly collected seeds was generally poor. Cold stratification (4°C) for 3 months in complete darkness improved germination, suggesting the dormancy breaking requirement in seeds.	Giménez-Benavides <i>et al.</i> (2005)
Empetrum nigrum	Seeds are dormant at the time of dispersal from plant and require warm stratification followed by cold stratification for germination. The seeds can also induce a secondary dormancy if the climate during the growing season is unfavorable.	Graae <i>et al.</i> (2008) and Baskin <i>et al.</i> (2002)
Stachys germanica L. subsp. bithynica	Freshly collected seeds did not germinate. Dormancy breaking requirements to break PD are satisfied by overwintering in soil surface (but GA_3 and Kinetin break dormancy in the laboratory). Thus the species persist in soil until spring and germinate readily once dormancy in broken.	Güleryüz et al. (2011)
Vicia sativa, V. angustifolia, V. amoena and V. unijuga	Seeds dispersed in autumn have PD or PY or combined (PY +PD) and require overwintering to break dormancy. Once dormancy is broken seeds germinate in spring.	Hu et al. (2013)
Euphrasia minima and E. salisburgensis	Although seeds can germinate at low temperature (5°C constant temperature or 3-10°C varying temperature) at the time of collection, after-ripening widens the temperature at which seeds germinate. Thus, germination synchronizes	Liebst and Schnelle (2008)

	with spring. This feature spread the germination over 3 years, thus these seeds formed a persistent seed bank.	
Silene elisabethae	Increase in duration of cold stratification increased the germination percentage indicating the cold stratification requirement to alleviate the non-deep PD. This requirement for dormancy alleviation should happen through overwintering in nature (but GA_3 in laboratory) stimulating germination in spring.	Mondoni et al. (2009)
Polygonum cuspidatum and P. weyrichii var. alpinum	Only moist chilling (cold-wet stratification was effective in breaking dormancy of high-altitude adapted <i>P.weyrichii var. alpinum</i> . Seeds germinate at 5°C following the dormancy breaking treatment, compared to 35°C requirement at the time of collection in autumn.	Nishitani and Masuzawa (1996)
Peucedanum ostruthium	After-ripening requirement for breaking (non-deep)-morphophysiological dormancy and high temperature requirement (26°C) delayed germination until spring.	Novak et al. (2011)
Achillea moschata, Hieracium staticifolium, Oxyria digyna and Rumex scutatus	Seeds sown in autumn only germinated in the spring suggesting cold stratification is required for germination in these species.	Stöcklin and Bäumler (1996)
Anemone narcissiflora var. sachalinensis, Arnica unalaschkensis, Bryanthus gmelinii, Bupleurum ranunculoides var. triradiatum, Carex flavocuspis, C. pyrenaica, C. stenantha var. taisetsuensis, Diapensia lapponica var. obovata, Empetrum nigrum var. japonicum, Fauria crista-galli ssp. japonica, Gentiana nipponica, Loiseleuria procumbens, Patrinia sibirica, Pennelianthus frutescens, Peucedanum multivittatum, Phyllodoce aleutica, Potentilla matsumurae, Primula cuneifolia, Rhododendron aureum, Sieversia pentapetala, Solidago virga-aurea ssp. leiocarpa, Therorhodion camtschaticum, Tilingia ajanensis, Vaccinium ovalifolium, V. uliginosum, V. vitis-idaea and Veronica stelleri var. longistyla.	Seeds of <i>E.nigrum</i> , <i>A. narcissiflora</i> and <i>G. nipponica</i> did not germinate at all irrespective of treatment. However, now it is now known that <i>E.nigrum</i> needs additional dormancy breaking treatment. Similarly, with the exception of <i>D. lapponica</i> , <i>V. uliginosum</i> , <i>P. frutescens</i> , <i>P. cuneifolia</i> and <i>C. flavocuspis</i> , all other species germinated to higher percentage under warm temperatures (35/25°C or 30/20°C). Cold-wet stratification given at 0°C for 2 months significantly reduced the temperature required for germination and cumulative germination percentage at all temperature regimes tested, i.e. 35/25°C, 30/20°C, 25/15°C, 20/10°C and 15/5°C.	Shimono and Kudo (2005)
Caltha leptosepala	At the time of dispersal, seeds are MPD. Dormancy is alleviated by stratification at 2.5°C, embryo growth begins within 1 month and germination begins in 4 to 7 months of storage. PD determines the germination timing.	Forbis and Diggle (2001)

II. Temperature requirement for germination

Most of the seeds endemic to alpine environments do not germinate immediately after shedding because the temperature required for germination is high (Sayers and Ward, 1966; Billings and Mooney, 1968; Densmore and Zasada, 1983; Nishitani and Masuzawa, 1996; Baskin and Baskin, 1998; Cavieres and Arroyo, 2001; Shimono and Kudo, 2003) and in autumn this temperature is rarely - if ever - met. After overwintering, cold-wet stratification reduces the high temperature requirement and seeds develop a tendency to germinate at a wide range of temperatures (Baskin and Baskin, 1998; Table 1).

It is not uncommon for some alpine species to shed nondormant seed that remain ungerminated during autumn due to the prevailing low temperatures (Billings and Mooney, 1968). Seeds of *Oxyria digyna* can germinate immediately after dispersal if the temperature is between 10 and 15°C, but some seeds can even germinate below 10°C, if the temperatures fluctuate between 13 and 2°C and have light for 12 h (Mooney and Billings, 1961). Neither cold-wet stratification, nor after-ripening is a strict requirement for germination, rather the germination after seed shedding was impaired by prevailing low temperatures in autumn. In accordance with this view, a warmer autumn temperature is likely to increase the germination of some nondormant seeds, e.g. *Geum reptans, Veronica alpina* and *Oxyria digyna* to autumn (Mondoni *et al.*, 2012). Such seeds are poorly represented in alpine communities, as the winter temperatures decimate seedlings (but see below).

Typically, 25 to 35°C is the most favorable temperature eliciting autumn germination in the proportion of non-dormant/conditionally dormant seeds (Clebsch and Billings, 1976; Kibe and Masuzawa, 1994; Nishitani and Masuzawa, 1996; Körner, 1999; Novak *et al.*, 2011). Autumn germination is facilitated further if the temperature fluctuates between 20 and 30°C (Sayers and Ward, 1966; McDonough, 1970). However, these relatively high temperatures do not occur in alpine environments during autumn (Billings, 1974; Bell and Bliss, 1980; Reynolds, 1984). Elkington (1971) showed freshly collected seeds of *Dryas octopetala* germinated to 48% when sown at 25°C under controlled germination conditions, but field sown seeds in autumn germinated only in the following spring, suggesting the temperature required for germination was not met. Likewise, Marchand and Roach (1980) also reported seeds of *Arenaria groenlandica, Juncus trifidus* and *Potentilla tridentata* germinated to 88, 88 and 52% when tested in laboratory at 23 to 28°C, 19 to 23°C and 21 to 26°C respectively, however, only 38, 17 and 6% of the seeds germinated in the field.

The mean autumn, spring and summer temperatures decrease with altitude. In addition, snow cover often comes early but lasts longer at higher elevations (Lütz, 2011). Evidence is mounting that the autumn and spring temperature optima for germination is different for species occurring at different altitudes. In general, seeds from higher altitudes where temperature minima is low have a tendency to germinate at lower temperature after cold stratification, but seeds from lower altitudes experiencing warmer conditions during seed development require higher temperature than high-altitude species (Dorne, 1981; Mariko *et al.*, 1993; Nishitani and Masuzawa, 1996). Thus, seeds moved from a low altitude in a reciprocal transplant experiment germinated in the autumn at higher elevations, but when seeds from higher altitudes were moved to lower altitudes seeds germinated only in the second spring following transplant (Jaganathan *et al.*, unpublished).

A simple germination experiment using *Polygonum weyrichii var. alpinum* and *P. cuspidatum* seeds at a range of temperatures including 5, 10, 20, 25 and 35°C, showed that

germination of seeds of both species could only be achieved in the autumn at temperatures of 35°C both in light and dark conditions (Nishitani and Masuzawa, 1996). Importantly, after cold-wet stratification, seeds of *P. cuspidatum* from Mt. Fuji (a site experiencing relatively high temperatures) germinated equally well at 10, 20, 25°C, contrasting with the poor germination of seeds from Shizuoka (relatively moderate temperatures) at 10°C. Similarly, in the case of *Phacelia secunda*, freshly collected seeds did not germinate under simulated diurnal fluctuations at 20/10°C or at 10/5°C at the time of collection in autumn, but seed germination increased proportionately with cold wet stratification duration (Cavieres and Arroyo, 2000). In particular, seeds from low altitudes required lesser stratification duration and germinated equally well at both the temperatures tested than did seeds from higher elevations (Cavieres and Arroyo, 2000). In another study, Blionis and Vokou (2005) confirmed the ability of higher altitude seeds to germinate at low temperature after overwintering in *Campanula* populations. Also, Mondoni *et al.* (2009) showed seeds of *Silene elisabethae* collected from the coldest site in their investigation germinated better at low temperature after cold-wet stratification.

Vera (1997) studying the effect of altitude and seed size on germination and seedling establishment of *Calluna vulgaris*, *Erica cinerea* and *E. vagans* collected from heathlands in northern Spain found that small sized seeds originating at higher altitudes germinated well at 20°C. A more recent study by Liu *et al.* (2013) on seed germination of 445 species in a grassland community on the eastern Tibet Plateau, demonstrated that after-ripened seeds germinate better at alternating temperatures (either at 5/25°C or 10/20°C) than at a constant temperature of 15°C especially if they were small forbs. More studies are needed to offer clearer understanding of autumn and spring germination ecology especially at the community level.

The local microclimate in fellfield environments is harsher than that of snowbeds due to little snow accumulation, and reduced snow cover thickness and duration (Körner, 2003). Snow patches covering soil for a longer time in snowbed fields offer refuge to the seeds from low temperature. For instance, seeds under snow cover are held in stasis and showed significantly lower level of damage and mortality than seeds kept artificially snow free (Williams, 1987). On average, growing season length decreases by 30-50% from fell-field to snow-bed. Thus, habit-specific temperature requirement for autumn germination is expected but there is no evidence for this trend. For example, seeds of *Potentilla matsumurae* collected from both fellfield and snowbed failed to germinate at the time of collection at 15/5°C, but with increasingly warm temperature cycles (20/10°C, 25/15°C, 30/20°C, 35/25°C) the germination percentage increased (Shimono and Kudo, 2003).

In the same study, Shimono and Kudo (2005) disclosed that seeds of some species e.g. *Arnica unalaschkensis*, *Bupleurum ranunculoides*, *Vaccinium ovalifolium* at the time of dispersal in autumn have a more moderate optimum temperature regime, thus germination at higher temperatures of 35/25°C is unfavorable. However, in these species cold-wet stratification has widened the temperature at which germination occurs. Overall, it seems logical to conclude that the temperature requirement for germination both in autumn and after overwintering in field is highly species-specific. Despite this, several sound pieces of evidence indicate that it is safe to generalize that the requirement for high temperatures for germination in autumn in alpine species can be beneficial to postpone germination to spring, especially in non-dormant seeds. Furthermore, studies that document germination temperature requirements have revealed both inter and intra-species variation in autumn germinating temperature, but acknowledge that the temperatures required for germination are not likely to be met in autumn.

III. Light requirement

Light is an important environmental factor stimulating germination in many seeds (Milberg *et al.*, 2000). The requirement of light for germination has been one of the potential determinants driving species to accumulate a persistent seed bank (Baskin and Baskin, 1998; Milberg *et al.*, 2000; Pons and Fenner, 2000). A strict light requirement in some alpine species not only prevents germination in late autumn, it also cues the timing of germination to spring (Densmore, 1997; Fig 1).

Seasonally changing day length acts as a bottle neck in controlling alpine seed germination. In Alaska, nonstratified seeds of Diapensia lapponica, Chamaedaphne calyculata, Saxifraga tricuspidata and Ledum decumbens showed no or few germinating seeds under short-day conditions (Densmore, 1997). However, cold stratification increased the proportion of germinated seeds of all species, significantly both under short-day and long-day conditions. Similarly, longer-day conditions promoted germination in all PD seeds tested, except Poa alpina (see below; Schwienbacher et al., 2011). Haggas et al. (1987) reported that seeds of Carex paysonis collected in Beartooth Mountain achieved good germination only under complete darkness followed by light conditions at constant temperature (25°C). Seeds incubated at 25°C under complete light or darkness resulted in poor germination. In addition to being dormant, seeds of C. pendula required light for germination even after the dormancy is broken (Brändel and Schütz, 2005). Seeds of Silene elisabethae, showed increased germination percentage with increase in cold-wet stratification duration, but germination was tightly controlled by light (Mondoni et al., 2009). Novak et al. (2011) also noted seeds of Peucedanum ostruthium are better germinated at 26°C only in light. These studies clearly depict light plays a pivotal role in controlling the timing of seed germination form seed banks.

By contrast, for some alpine species light is not required for germination, instead light inhibits it. Examples of dark-requiring species include *Poa alpina* (Acharya, 1989; Schwienbacher *et al.*, 2011), *Trifolium pallescens* (Schwienbacher *et al.*, 2011), *Anthyllis vulneraria* subsp. *alpicola* (Schwienbacher *et al.*, 2011), *Agropyron latiglume* (Acharya, 1989), and *Koenigia islandica* (Heide and Gauslaa, 1999; Wagner and Simons, 2009). Both dark and light requirement in seeds can play a pivotal role in delaying germination, thus contributing to seed persistence (Fenner and Thompson, 2005; Bewley *et al.*, 2013). Therefore, it seems reasonable to suppose that dormancy and light requirement both circumvent autumn germination in alpine species, thus affecting the composition of soil seed banks. In dark-requiring species (e.g. *Trifolium pallescens*), dormancy can act as the main mechanism in preventing seed germination until the growing season (Table 1). However, in spring, seeds experience light/dark cycles, therefore it is not known if cold-stratification in winter imposes a light requirement in the otherwise dark-requiring seed.

Important results from recent ecological studies suggest that environmental conditions significantly contribute to the development of a light requirement in alpine species. Seeds of *Koenigia islandica* collected from Iqaluit (Nunavut, Canada), Yukon (Canada), and Jasper (Alberta, Canada) did not have a significant germination light-requirement, whereas seeds collected from more severe climates, Svalbard (Norway) and Colorado (USA), required light for germination (Wagner and Simons, 2009). Mondoni *et al.* (2012) showed that seeds of *Oxyria digyna* germinated at 2°C higher than the current alpine climate (12/12 h photoperiod)

achieved higher germination, but that there is a tendency for the seeds to become autumn germinators because dormancy is lacking. Unfortunately, the role of light in germination control was not considered in their study. However, Billings and Mooney (1968) worked with *O. digyna seeds* and reported that those collected from an American alpine location had greater cumulative germination in light than in dark, but seeds of arctic origin did not have strict requirement for light. The aforementioned studies suggest the evolution of a requirement for light is more likely location-specific.

Light requirement in seeds has also been reported to be of great significance in controlling the depth of burial at which germination occurs (Pons and Fenner, 2000). Since light can only penetrate a few millimeters from the top of the soil surface, seeds found below this depth do not germinate; and even in dark-requiring species, those seedlings would die before reaching the soil surface, especially if they are small owing to the smaller seedling size (see below). Data on other ecosystems suggest that seeds that are smaller are more likely to require light for germination than larger seeds (Milberg *et al.*, 2000; Jankowska-Blaszczuk and Daws, 2007). In a recent study, Wu *et al.* (2013) found a similar pattern in alpine species of Qinghai-Tibetan Plateau. Using twenty species, those authors reported significant negative correlation between strength of light required for germination and seed mass, thus large-seeded species were less likely to require light for germination than those of small-seeded species. If small seeds are buried to greater depths, the strong requirement for light in small-seeded species suggests that disturbance is an inevitable requirement in the germination of seeds from seed banks. Nothing, however, is known of the light requirement of seed in different stages of the seed bank cycle (Fig. 1).

While the studies illustrated above lend some support to the assertion that light plays an important role in controlling seed persistence, this perspective requires detailed experimental investigations. Because our knowledge on the relationship between light requirement and germination is limited merely to genus *Carex* (Amen and Bonde, 1964; Schütz and Rave, 1999; Schütz, 2002) and *Koenigia islandica* (Heide and Gauslaa, 1999; Wagner and Simons, 2008, 2009), the understanding of the role of light in controlling the autumn germination of some non-dormant alpine seeds is still rudimentary. Nevertheless, we surmise that the long-day requirement for germination might be common (but not ubiquitous) in alpine species. This is because, in any year with an unfavorable growing season, long-day conditions will not be repeated until the following year, a condition typical to all alpine environments. However, in favorable years, a significant amount of light reaches the soil surface due to the lack of taller plants (Billings, 1974; Densmore, 1997). Hence, in addition to dormancy, light requirements also hold germination in check, thus contributing to seed persistence in soil.

IV. Factors affecting seed persistence

In many alpine species, the evolution of highly specific germination requirements, such as those described above, delay the germination of autumn-dispersed seeds until spring. Thus, the interplay of factors including dormancy, specific temperature requirement to germinate and highly specific light or dark requirements in a particular species offers triple safety measures against the harsh alpine climate; without these controls seeds are more likely to have asynchronous germination thereby posing a serious risk of seedling mortality (Billings and Bliss, 1959). For example, when seeds develop in a relatively warm climate due to interannual variation, the degree of dormancy (*sensu* Baskin and Baskin, 1998) may be reduced. In such cases, autumn germination may be prevented by a high temperature requirement and/or light requirement instead of the depth of dormancy acting as the primary

obstacle to germination. If only one mechanism for preventing germination has been evolved in alpine seeds, then the autumn germination ability is likely to depend on the maternal environmental condition during seed development and opportunistic temperature at autumn. Evolution of such complex germination requirements is not unique to all ecosystems around the world. The obvious example are flood meadows, where many seeds are non-dormant but germination is controlled by light and a narrow temperature requirement, hence dormancy has no role in seed persistence (Hölzel and Otte, 2004 and references therein).

In addition to a complex suite of responses to varying environmental cues, it has been suggested that not all the seeds of a particular species may respond to germination cues stimulated by seasonal temperature cycles in alpine environments, thus carryover of a proportion of seeds to next year is expected to maximize species fitness (Meyer and Kitchen, 1994a; Meyer et al., 1995). Carry over mechanisms are common in unpredictable conditions because the risk of germination is spread to different years. For some alpine species, e.g. Carex frigida, low germination in the growing season tends to favor carry over mechanisms (Schütz, 2002). Although the success of seeds carried over to germinate in the next year depends on various phenotypic and genotypic factors (Meyer and Kitchen, 1994a), how a species selects potential candidates for carry over is not understood completely. Thus, it is not known if the seeds to be carried over were predetermined at the time of shedding or the selection is a result of phenotypic variability or simply that all the non-germinated seeds at the end of growing season are carried over (but see Meyer and Kitchen, 1994a). However, the degree of dormancy, position of seeds in soil and local microclimate are certainly important factors determining which seeds are carried over. Furthermore, for various reasons that are typical to alpine environments, e.g. frequent summer frost (Marcante et al., 2012), only a proportion of seeds germinate even in favorable growing seasons (Cavieres and Arroyo, 2001), leaving the reminder to be part of the persistent seed bank.

(1) Seed size and persistence

A commonly accepted prerequisite for seeds to enter persistent seed banks is the small size and round shape (Thompson *et al.*, 1993). Despite their suggestion that this phenomenon is likely to be universal, subsequent studies testing the prevalence of small, round seeds in persistent seed banks have argued both for (Bekker *et al.*, 1998; Thompson *et al.*, 2001; Zhao *et al.*, 2011), and against (Leishman and Westoby, 1998; Moles *et al.*, 2000; Yu *et al.*, 2007) the hypothesis. There have been few works that studied the significance of seed size in establishing persistent seed bank in alpine landscapes. In Argentinean mountain grassland, the available evidence demonstrated that small seeded species have an increased likelihood of entering the persistent seed bank (Funes *et al.*, 1999; Fig. 2). Conversely, Cerabolini *et al.* (2003) showed a negative correlation between seed size and persistence in species tested on Italian Alps (Fig. 2).

Recent years have witnessed a surge of studies determining seed longevity in soil using artificial burial experiments. Schwienbacher *et al.* (2010) conducted such an experiment in the Central European Alps (ca. 3500m a.s.l; Austria) for *Achillea moschata*, *Artemisia genipi*, *Anthyllis vulneraria* sspp. *alpicola*, *Geum reptans*, *Linaria alpina*, *Oxyria digyna*, *Saxifraga aizoides*, *S. oppositifolia* and *Trifolium pallescens*. Seed germination following excavation at 1, 2 and 5 years revealed, with exception of *Geum reptans* and *S. oppsitifolia* (which have extremely tiny seeds), at least some of the seeds buried at a depth of 3 cm survived for a minimum of 2 years (Fig. 1; Table 2). However, their study had two surprising results. Firstly, in contrast to the suggestion that round seeded species have increased tendency to enter the persistent seed bank, flattened *L. alpina* seeds survived for 5 years forming a long-

term persistent bank, whereas other flattened seeds of *O. digyna* did not survive for 5 years. Secondly, species classified as transient by Cerabolini *et al* (2003), were shown to form short-term persistent or long-term persistent seed banks (Table1). The only other attempt to use an artificial burial approach to test seed viability for >1 year in alpine tundra was a study in the Chilean Alps (Arroyo *et al.*, 2004). From a total of 15 species studied, all but two species namely *Senecio magellanicus* (with cylindrical shape but smaller size) and *Berberis buxifolia* (6.33 mg), survived for two years. However, they terminated the experiment in two years, therefore a distinction between short-term and long-term seed persistence is not feasible.

There are several lines of evidence showing that small seeds easily become buried, thereby escaping post-dispersal predation, remaining in the same environment in which they had developed and becoming part of the persistent seed bank (Peco *et al.*, 2003; Ma *et al.*, 2010b). If this holds true then seeds at depth in the soil are expected to be smaller than the seeds in the surface layer. However, there are only a few experimental studies addressing the vertical movement of seeds in alpine flora. In the Austrian Alps, Klug-Pümpel and Scharfetter-Lehrl (2008) found small seeded species move to greater depths than large seeded species, but Ma *et al.* (2010b) failed to find such pattern in 122 species of Tibetan Plateau, China. The important question perhaps then is: what depth is safe for seeds? Ma *et al.* (2010b) showed that seeds buried (naturally) to 15 cm can germinate when excavated and sown in optimal germinating conditions. Seeds artificially buried to 3 cm deep germinated after 5 years (Schwienbacher *et al.*, 2010; Table 1). However, over the same time period but at 8-10 cm depth, Adzhiev *et al.* (2011) reported no germination in 45 out of 63 species of the Northwest Caucasus.

It is difficult to draw any firm conclusion on the impacts of seed size on persistence in alpine seed banks from these studies mainly because of the overriding effect of other factors influencing results, three of which are discussed here. The first to be considered is the type of soil surface where seeds land. If seeds are deposited on a surface having smaller particles, there is an increased likelihood that small seeded species rather than those with large seeds, and those with adhesive seed coats are trapped amongst the soil particles and buried in the soil (Chambers *et al.*, 1991; Chambers, 1995, 2000). Small soil particle size may enable the large seeded species to move horizontally over longer distances until they encounter soils with large particle sizes. The smaller seeds frequently trapped by soil of smaller particle size often move to a deeper position than the larger seeds trapped in large particle soil. In contrast, both small and large seeded species are buried easily if seeds were deposited on soil surface having larger particle size (see Chambers *et al.*, 1991 for a detailed discussion).

A second confounding factor when trying to generalize on the relationship between seed size and persistence is derived from the understanding that entry to the soil seed bank confers longevity. It is perhaps an over-simplification to say that ease of burial improves the longevity of small seeded species in the alpine system. Indeed, seeds buried to greater depths do not germinate, unless the soil is overturned but, if disturbance is very infrequent, these species may be lost from the aboveground community and eventually from the seed bank also. For example, in the central Chilean Andes, Arroyo *et al.* (1999) found buried seeds of *Chaetanthera pusilla* were viable, although the species did not occur in the standing vegetation. Similar results were obtained in the alpine regions of Tibet for a wide range of species: a total of 87 species were identified in the soil seed banks, but only 62 appeared in the standing vegetation (Ma *et al.*, 2010a). There are no data available on how long these seeds can be viable but presumably, this longevity will be species-specific. Nonetheless, dating by accelerator mass spectrometry revealed seeds of *Carex bigelowii* and *Luzula*

parviflora in Alaskan tundra had at least persisted for two to three centuries (McGraw et al., 1991).

A third problem with linking seed size and persistence is the inconsistency in methods with which seed longevity is determined; this causes subsequent confusion when trying to build causal relationships between size and viability. For example, seeds of some species classified as transient by Cerabolini et al. (2003), survived more than 1 year when tested with artificial burial experiments (Table 2). It has been suggested that the discrepancy in the findings of these studies has resulted from different methods employed in testing seed longevity (Schwienbacher et al., 2010). Cerabolini et al. (2003) used published information and germination experiments conducted with seeds excavated from soil kept in glass houses to classify the seed bank type, whereas Schwienbacher et al. (2010) classified the seed bank type based on data from artificial burial experiment. Each method has its own advantages and disadvantages; the artificial burial approach often overlooks the mechanism of burial, which is arguably a complex process itself (Thompson et al., 1997). Therefore, the conclusion of Schwienbacher et al. (2010) that Linaria alpina retains its viability in the soil for 5 years as estimated by artificial burial experiments, can be called into question because the soil environment may not enable adequate burial and subsequent avoidance of disturbance to maintain the conditions generated in their experiment. Thus, whether shape is a good predictor for seed persistence or not, remains largely equivocal for alpine species. More meaningful answers to this question will only come from novel seed bank studies that combine both natural soil sampling and artificial burial approach incorporating small and large seeds.

While some small seeded species more easily enter persistent seed banks than large seeded species, careful consideration of seed persistence studies point to the possibility that not all small seeded species can potentially establish persistent seed banks (Fig. 2). Similarly, generalizations on large seeded species are inappropriate as some of these can also establish short-term persistent seed banks (Fig. 2). As yet, there is no evidence available to demonstrate whether large seeded species possess a long-term persistent seed bank in alpine landscapes (Fig. 2). However, a short-term persistent seed bank tends to be adequate for maintaining species in alpine conditions, because the favorable weather conditions can occur at least once every few years (Körner, 2003). Furthermore, most of the plants adapted to alpine tundra are perennials and produce seeds in fairly large amount, at least in the favorable years, and although a long-term persistent seed bank is adaptive to the alpine climate, it is not necessary for a species to succeed in colonization.

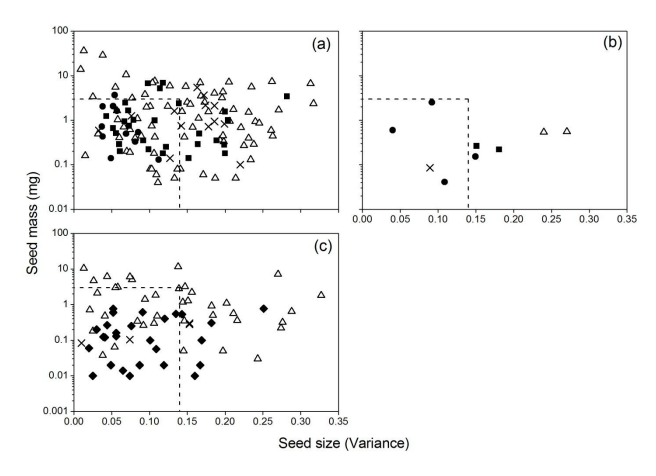


Fig. 2 Relationship between seed mass (log scale) in mg and variance of seed dimension adopted from (a) Italian Alps; 114 species (re-drawn only for alpine species from Cerabolini *et al.*, 2003 with permission), (b) Central Austrian Alps (Schwienbacher *et al.*, 2010) after excluding the standard deviation given in the original data set and (c) Argentinean mountain grassland; 71 species (re-drawn from Funes *et al.*, 1999 with permission). (Δ) Transient, (•) Long-term persistence, (■) short-term persistence, and (•) persistence (but no information on whether short or long-term), (X) uncertain or could not classified. More detailed information of how to calculate the variance of dimension can be found in Thompson *et al* (1993). In order to make some extremely small seeded data point readable in the Argentinean mountain grassland (c) we have rescaled the x-axis of this study to begin from 0.001.

(2) Seedling establishment

An additional factor that complicates our understanding of the significance of seed size in persistence is the hypothesis related to seedling establishment. With finite resources allotted to reproductive success, a given species may produce more small seeds or less large seeds, leading to a variation in seed mass often spanning five to six orders of magnitude across species found in a particular ecosystem (Thompson, 1987; Jakobsson and Eriksson, 2000; Leishman *et al.*, 2000; Leishman and Murray, 2001). A growing body of literature is specifically devoted to understand how such variation in seed size influences the fitness of plant communities (Leishman and Murray, 2001). Although small-seeded species benefit by producing more number of seeds than large seeded-species and have an increased likelihood of entering soil banks, the latter group might counteract the advantage by establishing high number of successful seedlings than the small seeded-species.

Because seed germination is an irreversible event, timing of germination is a critical component in the life-cycle of alpine plant species. Seeds, once germinated, have to continue growing or die. Seedling establishment in alpine environments is a slow process and often lasts for several years (Jaganathan, personal observation). Although some of the seeds germinating under snow cover are very slow requiring up to 6 months to achieve 50% germination (Meyer *et al.*, 1995), spring germinators must attain a suitable size to tolerate subsequent winter conditions (see Leishman *et al.*, 2000), if not, seedling mortality becomes prevalent. If this holds true, then it is reasonable to expect that large-seeded species establish seedlings more quickly and there may be a trade-off between seed size and mean germination time (MGT). Consequently, we anticipate that large-seeded species must germinate faster than small ones to maintain an advantage over the seed bank strategy prevalent in small-seeded species.

We tested this hypothesis in 653 Qinghai-Tibet Plateau species (original data from Bu *et al.*, 2007a; Wu *et al.*, 2013). Species were only included in this analysis if they were recorded as germinated. Hence, species recorded with MGT of 0 (failed germination) were excluded from this analysis. In this way, we were able to select a total of 585 species from 39 families. We performed a linear regression to explain the relationship between seed mass and their respective MGT. In contrast to our expectation, we observed no significant relationship between seed mass and MGT (slope = 1.8, $R^2 = 0.01$, d.f. = 584, p > 0.05) (Fig. 3).

If large seed size does not confer an advantage with respect to MGT an alternative benefit may be bestowed through greater seedling size (Vera, 1997; Moles and Westoby, 2004). However, dormancy can regulate germination only in terms of when in the growing season emergence occurs for more successful seedling establishment. Further, there is little evidence that large-seeded species have some advantage in survival over small-seeded species (even within same genus, see Veera, 1997). We conclude that this advantage may not counterbalance the large number of seeds produced by small-seeded species and their ability to persist in soil.

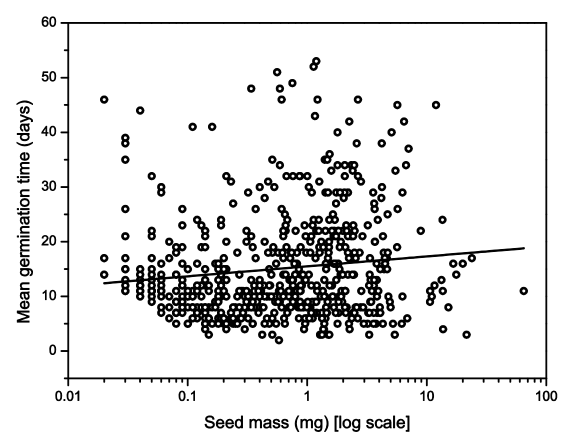


Fig. 3 The relationship between seed mass and MGT (Mean Germination Time) estimated for 565 Qinghai-Tibet Plateau species. Number of days required for germination increased with increase in seed size (slope =1.8) but the relationship is not significant ($R^2 = 0.01$, d.f. = 584, p > 0.05). Each point represents one species.

(3) **Seed dispersal**

At the ecosystem level, microhabitat and seed availability determine the composition and diversity of species (Bullock et al., 1995; Tackenberg et al., 2003; Ozinga et al., 2004). Seed dispersal has a marked effect on seed availability in any given location and the distribution of the 'seed rain' is potentially very different to that of the parent plants. Plants adapted to shortgrowing seasons in unpredictable environments are believed to disperse seeds over a longer distance, so that seeds experience different weather conditions and germinate in the environments where conditions are favorable (Bakker et al., 1996; Baskin and Baskin, 1998; Tackenberg et al., 2003). Given the unpredictability in climate, it is reasonable to expect alpine species have long distance dispersal (LDD). In spite of a pullulating interest on understanding the seed dispersal in low-altitude ecosystems, our knowledge on seeddispersion patterns for alpine ecosystem remains conjectural (McGraw and Vavrek, 1989; Muñoz and Arroyo, 2002). In particular, research on alpine seed dispersal has mostly concerned the mode of seed movement, rather than precisely estimating how long seeds could move. One reason for this lacuna is the difficulty in tracking all routes a seed could move from mother plant to the germination site. Even for the studies that attempted to quantify the distance, results were influenced by several other factors such as height from which seeds are dispersed; size and morphology of seeds; morphological adaptations for dispersal; dispersal vectors availability; wind speed at the time of dispersal and wind direction.

Seed movement within alpine landscapes involves numerous dispersal vectors. Most frequently seeds are dispersed by wind (Van der Pijl, 1982; Willson *et al.*, 1990; Table 3). A simulation model in the glacier foreland of the Dischma Valley, Switzerland has shown that more than half of the alpine species were dispersed a long distance by wind, whereas only 25% of low altitude species tend to have LDD (Tackenberg and Stöcklin, 2008). Because the horizontal movement of seed can be affected by the wind speed, Tackenberg & Stöcklin (2008) also measured the wind speed in alpine and low-altitude, but found only a small difference between these two environments. However, wind speed may not only vary between season but also years (Billings and Bliss, 1959). Furthermore, secondary movement of seeds also depends on the soil characteristics (Chambers *et al.*, 1991; Chambers, 1995, 2000). The small-sized seeds frequently trapped by smaller particle size move vertically, thus do not move to long-distances (see Chambers *et al.*, 1991 for a detailed discussion). Therefore, it is not known if the simulation model has overestimated the role of soil characteristics and wind speed in secondary seed dispersal.

The dogma that seed dispersal agents are rare in alpine environments (McGraw and Vavrek, 1989), has been disproved by active research over the last two decades. Seeds of many alpine plants could successfully survive passage through the intestinal tract of animals, i.e. endozoochory (Bruun *et al.*, 2008; Yu *et al.*, 2012). For some species, this passage promoted germination, whereas for others it suppressed the total percentage of germinating seeds. A good example is the Andean (Chile) shrub *Berberis empertifolia* dispersed in the faecal deposition of *Liolaemus bellii*. After passing through the digestive tract, germination increased significantly when compared to the untreated control (Celedon-Neghme *et al.*, 2008).

Table 3 Distance seeds moved by wind.

Species	Distance moved	Reference
Ranunculus adoneus	30 cm	Scherff et al. (1994)
Arenaria groenlandica, Juncus trifidus, Potentilla tridentata and Diapensia lapponica	1m	Marchand and Roach (1980)
Erythronium grandiflorum	1m	Weiblen and Thomson (1995)
Campanula thyrsoides	<10 m	Frei et al. (2012)
Epilobium fleischeri, Senecio incanus, Genum reptans, Oxyria digyna, Saxifraga bryoides, Doronicum clusii, Linaria alpina, Poa alpina and Leucanthemopsis alpina	>100 m	Tackenberg & Stöcklin (2008)

In the alpine forest of southwestern China, species of *Pinus armandii*, *P. densata*, *Abies sp.* and *Viburnum sp.* were predated by rodents (Wang *et al.*, 2012). Two years of a seed caching experiment showed not only that seeds of *P. armandii* were dispersed to < 10 m but also rodent seed selection for caching was size-specific. In the experiments conducted with two populations of endangered Spanish black pine *P. nigra* in the Cuenca Mountains of Central Spain, seed predation in a mast year was significantly lower compared to the low seed producing year (Lucas-Borja *et al.*, 2012) suggesting seed predation varies between years. Mountain snowberries (*Gaultheria depressa*) endemic to New Zealand alpine ecosystems was consumed by scree weta (*Deinacrida connectens*) and moved long distances (Larsen and Burns, 2012), however such dispersal events are rare.

The ability of ants to disperse seeds was assessed experimentally in Central Chilean Andes for Sisyrinchium arenarium (Iridaceae) (Muñoz and Arroyo, 2002). Insightful measurements made at two different altitudes (2700m and 2000m) revealed greater activity of ants and birds at higher altitude compared to lower altitude, however, the overall seed removal by ants was significantly greater than birds at both elevations. In a subsequent study those authors showed that 9 species namely Alstroemeria pallida, Anarthrophyllum cumingii, Chuquiraga oppositifolia, Laretia acaulis, Rhodophiala rhodolirion, Sisyrinchium arenarium, Taraxacum officinale, Azorella monantha and Pozoa coriacea, were dispersed by ants and birds, but only over a short distance (Muñoz and Cavieres, 2006). In the sub-alpine locations of Mt. Norikura, Kita-Alps, Japan, seeds of Dicentra peregrina were observed to be dispersed by Formica gagatoides, but the distance is not known (Komatsu et al. 2014). To the best our knowledge, experimental evidence of ants dispersing seeds in other alpine environment has not been documented elsewhere. However, a growing number of studies are explicitly showing the significance of seed dispersal by birds in other alpine locations. For instance, in New Zealand, some fruit species were dispersed only by a fruit-eating parrot, kea (Nestor notabilis) (Young et al., 2012), but the specific distance of seed movement was not measured.

In New Zealand, 10 *Veronica* species disperse their seeds by the mechanism of hygrochasy (Garnock-Jones and Lloyd, 2004; Pufal *et al.*, 2010). Hygrochasy is generally understood as the ability of seed holding structures to dehisce when the moisture content of those structures increases to a certain threshold level. This mechanism of seed dispersal is common in desert species, where water availability and seed dispersal (and germination) must coincide. The evolution of hygrochasy beyond these alpine endemics needs further clarification, although it is assumed that this mode of dispersal does not favor LDD.

Many alpine species were able to float in water for more than one year, when empirically tested (Danvind and Nilsson, 1997). Nevertheless, long-distance movement of seeds by floating may be constrained by several factors (Danvind and Nilsson, 1997). Since alpine landscapes are covered by snow for most of the year, there is a greater probability that the seeds dispersed outside the snow-free growing season land on the snow blanket. Although these seeds are dispersed by wind (Matlack, 1989; Chambers and MacMahon, 1994), accumulating snow cover precludes further movement once the seed has come into contact with the snow surface. During snow melting these seeds move, but only over a short distance. For example, the work by Scherff *et al.* (1994) showed melting snow moved seeds of *Ranunculus adoneus* only to a distance of 10 cm. Similarly, Glaser (1981) observed extremely scarce movement of seeds when snowbeds melted in Mount McKinley National Park, Alaska. Evidence of this kind clearly demonstrates that the distance seeds move by this mode is inconsequential (Greene and Johnson, 1997).

In recent years, the influence of humans on alpine vegetation has become more pronounced. The construction of roads or walkways, regular visits to specific areas for research and education, and the recent surge of interest in mountain tourism frequently disturb vulnerable alpine communities. In addition, man and man-made machines are more likely to move lowland seeds in to alpine environments. Emerging studies have just started to understand the severity of the consequences for alpine plants, but for some lengthy treatment of this subject consult Pauli *et al.* (2001). Trampling damage due to walkers entering the Central Tasmanian alpine vegetation affects the shrubs, shrubland and grassland, although fen tend to be more resistant (Whinam and Chilcott, 1999). Non-native seeds belonging to 17 families have been identified in polar regions as a result of accidental transport by visitors (Ware et al., 2012). In Mount Norikura (Japan), construction of road has changed the species composition as a result of alteration in canopy and soil nutrition (Takahashi and Miyajima, 2010).

Except for wind dispersal, our knowledge of seed movement and potential dispersal distances by other modes remains far from satisfactory. Indirect evidence suggests that limited seed availability due to short-distance dispersal impedes the regeneration potential in open alpine landscapes rather than microhabitat availability. For example, restricted dispersal of *Vaccinium myrtillus, Rhinanthus minor, Rumex acetosa, Cicerbita alpina, Trollius europaeus* and *Campanula thyrsoides* appears to be the foremost reason for the patchy distribution, because seeds added manually to unoccupied sites resulted in successful germination (Lindgren *et al.*, 2007; Frei *et al.*, 2012).

It has been argued above that LDD is a sporadic event in alpine species. Whilst this is based on limited evidence, we nonetheless expect a larger set of data might yield similar results. As of now, the notion that LDD is rare in alpine species should be viewed as a working hypothesis rather than conclusion.

(4) Dispersal vs. persistence

Given finite reserves of energy to establish their off-spring successfully, many plants have evolved different ways of ensuring dispersal to different locations or persistence in soil to reduce the so-called 'colonizing effect'. One way is to invest in seed dispersal structures to attain longer dispersal. Information regarding the seed dispersal structures for alpine flora are incomplete, but a few alpine plants have been reported to invest in seed dispersal structures (Chambers and MacMahon, 1994). Bu *et al.* (2008) provided substantial data on the seed dispersal structure at community level in eastern Qinghai-Tibet plateau (Fig. 4). This data set suggests that most of the alpine plants present do not invest in any specific dispersal structures (Fig. 4), although a few species do invest in wind dispersal structures. In their work on comparing seed dispersal patterns for various plant communities, Willson *et al.* (1990) noted many alpine species could have morphological adaptations for wind, thus seed dispersal by wind may be common, amongst other available dispersal vectors.

While the studies by Willson *et al.* (1990) and Bu *et al.* (2008) generally agree that most alpine species neither invest in specialized morphological structure nor move long distances, the question as to why this is the case cannot be answered from available evidence. There may be several plausible explanations, although none of them have been rigorously treated in experimental works. Due to plant development in extremely cold-climates, the parent plant may have only limited resources allocated for seed development (Forbis, 2003; Körner, 2003). Larger seeds with specific dispersal structure means augmented investment, a reason that might explain why there were fewer fleshy seeded species observed by Willson *et al.* (1990). Furthermore, seed development must take place in a short time, if not, the

development is carried over to next year – principally because most alpine plants (and plant parts) are held in stasis by the snow cover except the seed producing and growing season.

A decade ago, Körner (2003) tentatively concluded that most of the alpine species may have seed mass of less than 4 mg. Extending his suggestion to a larger data set of more than 1000 species by reworking the data presented by Ma *et al.*(2010b), Funes *et al.* (1999), Cerabolini *et al.* (2003), Körner (2003), Giménez-Benavides *et al.* (2005), Pluees *et al.* (2005), Shimono and Kudo (2005), Bu *et al.* (2007a), Schwienbacher (2010), Ma *et al.* (Ma *et al.*, 2013) [from the seed size summary data] and Wu *et al.* (2013), we show that 88% of the species have seed mass below 3 mg and nearly 93% have seed mass below 4 mg, reinforcing the selection of small-seeded species in alpine vegetation. Furthermore, in literature where seed mass data are not presented for individual species, those studies have some useful summary of seed mass range. For example, of the 122 species reported in the study by Ma *et al.* (2010b), only few were above 3 mg, with the seed mass of 4.3618 mg (*Vicia multicaulis*) as the largest seed.

Baker (1972) first proposed that seed weight decreases with elevation using a data set from California. However, successive studies have either rejected (Mariko et al., 1993; Pluess et al., 2005) or accepted (Bu et al., 2007b; Guo et al., 2010; Dainese and Sitzia, 2013) this proposition, thereby undermining the generality of this pattern. One plausible reason for lack of unanimity in results is because in these studies the climatic conditions in which the seeds have developed were poorly measured or ignored. It has been stated that the short length of summer and low temperatures at high-altitude potentially limits the seed development duration and 'there may simply be inadequate time available for producing and stocking large, heavy seeds' (Baker, 1972). To support this view, seed developmental studies of Gentianella germanica comparing histogenesis and maturation drying over three years, have indicated that seeds developed in colder years were smaller and poorly developed when compared with seeds from the warmer year (Wagner and Mitterhofer, 1998). Similar conclusions have been presented for numerous species including Potentilla pulcherrima (Stinson, 2004).

Another limitation of the debate on the impact of seed size is due to the differing objectives and methods used to determine if a relationship exists between seed size and elevation. For Baker (1972) the average seed weight of species occurring at a particular altitude compared with the mean seed weight at a different (higher) altitude shows a decreasing trend, although discussions on one (Penstemon) genera and possibly same results on other genera were presented. Pluess et al. (2005) measured the seed size in 29 Swiss alpine congeneric species and found seed size increased with altitude for 55% of the species tested, but decreased or remained equal for 3% and 41% of species respectively. Nevertheless, populations of Scabiosa lucida, Saxifraga oppostifolia, Epilobium fleischeri and Carex flaccafound occurring at different altitudes showed a decrease in seed size with altitude. Mariko et al. (1993) working with Reynoutria japonica seeds collected from various altitudes of Mt. Fuji, concluded that mean seed weight of the upland population was significantly heavier than that of the lowland populations. Interestingly, the proposition that seed size increases with altitude is attributed to the fact that seedlings from larger seeds are more likely to survive extreme conditions than seedlings developed from small seeds. However, if seedling size advantage is a reason for the increased seed size observed with altitudes, this does not explain why large seeded species are rare in high altitudes (Table 4). Although we are inclined to believe that having a larger seed in comparison to low altitude species may be advantageous for alpine species of higher altitudes, we also accept that this 'trend' is not universal - even within an individual plant the seeds produced vary in size significantly, presumably due to numerous intrinsic and extrinsic factors, e.g. maternal environment, heteromorphism etc.

Table 4 Seed size of 1011 species collected from the published literature, Ma *et al.*(2010b), Funes *et al.* (1999), Cerabolini *et al.* (2003), Körner (2003), Giménez-Benavides *et al.* (2005), Lorite *et al.* (2007), Pluees *et al.* (2005), Shimono and Kudo (2005), Bu *et al.* (2007a), Schwienbacher (2010), Ma *et al.* (Ma *et al.*, 2013) [from the seed size summary data] and Wu *et al.* (2013). For some species, seed mass was determined by searching the seed information database (Royal Botanic Gardens Kew Seed Information Database (SID))

Seed size (mg)	Number of species
0-3	886 (87.6 %)
3-6	82 (8.1 %)
6-9	20 (1.9 %)
9-12	9 (0.9 %)
12-15	5 (0.5 %)
15-18	3 (0.3 %)
18-21	1 (0.1 %)
21-24	2 (0.1 %)
27-30	1 (0.1 %)
36-39	1 (0.1 %)
63-66	1 (0.1 %)
Grand total	1011

The underlying hypothesis describing the evolution of alpine plants has not yet been completely understood. However, it is suggested that some of the spermatophytes colonized tundra vegetation by dispersing seeds with differing phenotypes, some of which could survive extremely complex and uncertain alpine weather (Billings and Mooney, 1968; Marchand and Roach, 1980). The exact empirical evidence verifying this hypothesis cannot be possibly secured, but this view appears true, especially in terms of recent literature contemplating the encroachment of low-altitude species into the alpine vicinity as a consequence of global warming providing optimal survival conditions (e.g. Pauli *et al.*, 1996; Gottfried *et al.*, 2001; Parolo and Rossi, 2008; Walther *et al.*, 2009).

Many theories have been proposed to explain the drivers behind evolution of seed selection and adaptation. Hubbell's (2001) neutral theory of biodiversity assumes all individuals of all species are functionally equivalent, thus species abundance in a community is driven by the stochastic events such as dispersal, local extinction and speciation. This is in contrast to niche theory, which claims each species has its own traits and these independent species co-exist within a community (Coomes *et al.*, 2002). Leishman and Murray (2001) argued that seeds of early successional species were often smaller with good dispersal capacities. Such seed traits were advantageous in evolution, as a higher number of seed increases the probability to find different climates and germinate on suitable microsite. However when succession proceeds, both small and large-seeded species become more common, possibly due to the ability to persist in soil seed banks (either influenced by environment or maternal affects). More recent studies have therefore rejected the prediction of neutral theory (McGill, 2003; Gilbert *et al.*, 2006), indirectly supporting the overriding role of environment in selecting the phenotype.

It has been shown in Qing-Hai Tibet Plateau that early successional species were small but achieved high abundances, thus, seed size acted as a main determinant of species abundance (Chu *et al.*, 2007). In addition, seed mass was negatively related to species density early in the succession and this relationship became insignificant with time, implying large-seeded species started colonizing in later successional stages. This idea has been reinforced by examining the community assembly changes in Qinhghai-Tibetan Plateau China, where abundant species in an early-successional meadow were shown to be small-seeded species (Zhang *et al.*, 2012). However, the question as to what extent environment acted as a selective pressure and determined the evolutionary origin of alpine seeds along the successional gradient still remains unclear.

These studies, although few in number, offer insights suggesting that, owing to the advantage that species with smaller seeds have in entering seed banks (Funes *et al.*, 1999; Cerabolini *et al.*, 2003; Schwienbacher *et al.*, 2010), most alpine species may benefit from the seed banks rather than dispersal. Furthermore, seeds in alpine soil remain frozen most of the year and therefore secondary dispersal could be limited during these periods. This feature essentially drives the seeds to establish a soil seed bank. Consequently, entry into a seed bank may be critically advantageous not only because little to no investment for separate structures is required but also because the low soil temperature ensuring increased longevity of seeds (McGraw *et al.*, 1991). We therefore suggest that most alpine seeds have an increased tendency to be found in a seed bank in the model developed by Chambers *et al.* (1994).

This review has thus far dealt with various factors that influence the entry and exit of seeds into alpine seed banks but all of these bar the external environment are strongly affected by the phylogeny of the species themselves. The relatedness of taxa in alpine systems therefore has the potential to explain some of the variation in seed size and morphology, dispersal strategy, dormancy mechanisms and response to germination cues, all of which affect the

composition of seed banks. Links between phylogeny and traits affecting seed bank membership have been made in temperate systems (Mazer 1989; Leishman *et al.*, 1995), tropical communities (Norden *et al.*, 2009) and alpine/subalpine vegetation (Bu *et al.*, 2007a; Bu *et al.*, 2008; Xu *et al.*, 2014; Zhang *et al.*, 2004). More generally, Finch-Savage and Leubner-Metzger (2006) highlight the common evolutionary path for dormancy that means it is present throughout the higher plant families with only a few exceptions, and in all major climate zones. However, associations between phylogenetic groupings and germinability (measured as germination percentage, germination time, or both) whilst being significant, are very variable and models point to a number of sometimes unidentified alternative explanators. For example, Bu *et al.*, (2008) could explain only 12% of the variation of germination time in a one way ANOVA by using phylogeny, whilst Bu *et al.*, (2007) attributed 65.4% of germinability variation to genus-level groupings using GLMs that also included habitat, a non-significant explanatory variable. In addition to these ambiguous results, there is a geographic focus to the plateau communities of central Asia and until researchers in other regions emulate their efforts, no generalities can be made.

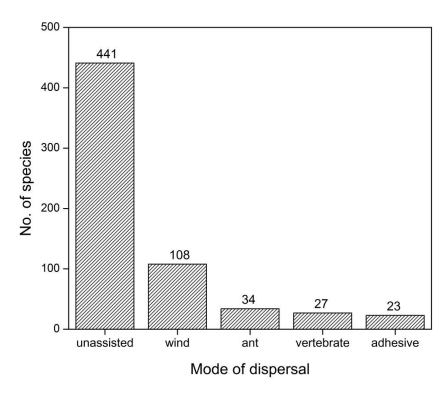


Fig. 4 Dataset of 633 species collated by Bu *et al.* (2008) showing number of species investing in specific dispersal structure.

V. Conclusions/future research needs

Understanding how alpine seeds establish seed banks is of extensive ecological and practical importance, because changes induced by global warming directly affect seed regeneration potential, thus influencing the floristic changes of alpine environments. Our ecological understanding of seed persistence in alpine ecosystems is still in its infancy, although seed persistence in soil seed banks is clearly an ecologically important trait for most alpine species. It is therefore not surprising that most of our discussion stems from studies addressing seed adaptation in alpine environment, rather than seed bank studies *per se*. Many areas have received only sporadic attention and data from those studies can be subjected to criticism for their disagreement in results and also due to bias in sampling. For example, a handful of studies (e.g. Schwienbacher *et al.*, 2011) have stored seeds at a condition that would benefit the physiological changes favoring dormancy alleviation and progress the seeds towards germination before being used in original experiments. Needless to say, such shortcomings must to be avoided in order to precisely understand the effects of maternal environments imposed on seed development. In addition, we have identified several areas that require future research with emphasis on the following:

- (1) The inherited traits of dormancy, germination temperature and light requirement could be affected to some extent by the maternal environment, which exert a selective pressure in developing physiological mechanisms that are location-specific. The effect of maternal climate affecting some of the adapted traits clearly suggests that these traits are likely to be altered in a future warming climate. Evidence for these patterns is already becoming apparent in some of the alpine locations. However, much remains to be studied.
- (2) It is suggested that climatic conditions in the growing season only partly controls seed carry-over. Because of specific germination temperatures required for each species present in a population, some temperatures preclude germination in some species and carry the seeds forward; the same temperature would initiate germination in other species. Since not all dormant and small seeded species enter persistence seed banks, there is a possibility that some other (physiological) factors are also involved in persistence under alpine climate. In addition to finding such factors, if any, contributing persistence, we need more rigorous appraisal of seed characteristics in relation to the environment. These approaches should also take *inter alia* inter-annual and inter-population variability into account. Thus, more specific studies delineating the point at which seeds bury in the model (Fig. 1) are needed.
- (3) Driving forces selecting seeds for seed banks must be evaluated in detail. Seed loss by germination and dispersal should be documented clearly. More especially, knowledge on dispersal is important in spatial demography studies particularly on soil seed banks. Many studies have shown vegetation of alpine regions is patchy possibly due to limited dispersal. LDD rarely occurs in alpine regions. If seeds do migrate to long-distance, habitat-dependent germination requirement are not met often, thereby seeds either germinate asynchronously to the dispersed environment or remain in soil for long time.
- (4) While a limited number of recent studies mostly on Qinghai-Tibet plateau, China, have focused on dispersal structure, little –if any- information exists on communities from other locations. This limited data on seed dispersal structure and dispersal distance dataset clearly underpins the need for more community level studies, paying particular attention to altitudinal and climatic influence.

- (5) Further phylogenetic investigations are needed to tease out the relationships between the key traits affecting the entrance to, and persistence in, the alpine seed bank and evolutionary origins of various responses to alpine environments. Furthermore, research in many more alpine regions other than central Asia is needed. Phylogenetic explanators of seed bank membership, if they exist, would allow more insightful predictions of responses to environmental change where species composition of the above- and below-ground communities is known, but responses of seed germination under warming conditions is untested.
- (6) A critical understanding of the germination requirements of local and invading species is essential. Many species probably survive changing climate by migrating to the locations where the originally adapted-for conditions still prevail (Hughes, 2000; Walther *et al.*, 2009). Climate change has now surpassed a threshold, inexorably forcing us to get a glimpse of types of plants that would survive in the future climate and in so doing, more meaningful conservation strategies can be devised. There is persuasive evidence that global warming will result in migration of low-altitude plants to alpine environments. The lower likelihood of alpine seeds reaching new environments highlight the need to recognize the interaction of encroached species with the endemic alpine seeds germinating from soil seed banks. In the future climate, these interactions essentially determine the species composition in alpine environment.

Insights into these areas will essentially help reveal the selective forces behind the seed adaptation/selection in alpine environment. Such results also would provide more understanding on how warming climate affects alpine seed banks.

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