UC Merced

UC Merced Previously Published Works

Title

The genetics of drought tolerance in conifers.

Permalink

https://escholarship.org/uc/item/2px0h6xh

Journal

The New phytologist, 216(4)

ISSN

0028-646X

Authors

Moran, Emily Lauder, Jeffrey Musser, Cameron et al.

Publication Date

2017-12-01

DOI

10.1111/nph.14774

Copyright Information

This work is made available under the terms of a Creative Commons Attribution-NonCommercial-NoDerivatives License, available at https://creativecommons.org/licenses/by-nc-nd/4.0/

Peer reviewed







Tansley review

The genetics of drought tolerance in conifers

Author for correspondence: Emily Moran Tel: +1 209 631 5194 Email: emoran5@ucmerced.edu

Received: 20 March 2017 Accepted: 27 July 2017

Emily Moran¹, Jeffrey Lauder¹, Cameron Musser², Angela Stathos¹ and Mengjun Shu¹

¹UC Merced, 5200 N Lake Rd, Merced, CA 95343, USA; ²Yale School of Forestry & Environmental Studies, 195 Prospect Street, New Haven, CT 06511, USA

Contents

	Summary	1	٧.	Implications for the management of forests in a changing world	9
l.	Introduction	1	VI.	Future directions	10
II.	Drought tolerance	2	VII.	Conclusion	12
III.	Investigation of the genetic basis of drought tolerance: three main approaches	4		References	12
IV.	Conifer drought tolerance genetics: what we know	5			

New Phytologist (2017) doi: 10.1111/nph.14774

Key words: association, conifer, drought tolerance, gene expression, genetics, genomic, local adaptation, provenance.

Summary

As temperatures warm and precipitation patterns shift as a result of climate change, interest in the identification of tree genotypes that will thrive under more arid conditions has grown. In this review, we discuss the multiple definitions of 'drought tolerance' and the biological processes involved in drought responses. We describe the three major approaches taken in the study of genetic variation in drought responses, the advantages and shortcomings of each, and what each of these approaches has revealed about the genetic basis of adaptation to drought in conifers. Finally, we discuss how a greater knowledge of the genetics of drought tolerance may aid forest management, and provide recommendations for how future studies may overcome the limitations of past approaches. In particular, we urge a more direct focus on survival, growth and the traits that directly predict them (rather than on proxies, such as water use efficiency), combining research approaches with complementary strengths and weaknesses, and the inclusion of a wider range of taxa and life stages.

I. Introduction

Climate change is increasing water stress in many areas as a result of increasing evaporative demand, altered precipitation and earlier snowmelt (Ryan, 2011; IPCC, 2013). Moderate drought is a common cause of reduced growth and increased mortality (van Mantgem & Stephenson, 2007; McDowell et al., 2010), whereas severe droughts have led to mass mortality (Breshears et al., 2005; Allen et al., 2010). Differences in drought susceptibility between tree species and populations have both environmental and genetic components. There is evidence of local adaptation to climate in many trees despite high gene flow (Aitken et al., 2008). Populations

often do not respond equally to a given climate (Rehfeldt et al., 2014). To better manage tree populations, we need to understand the relationships between tree genotype and phenotype (Gailing et al., 2009; Neale & Kremer, 2011) and between phenotype and ecological function (Tyree & Ewers, 1991; Sevanto et al., 2014; Montwé et al., 2015; Reinhardt et al., 2015). The relative importance of drought response plasticity vs genetic differences is still largely unknown, as are details of which traits are most important for drought tolerance and what genes underlie them.

Here, we focus on the genetics of drought tolerance in conifers. Although some general principles apply to both angiosperms and gymnosperms, there are significant differences. Gymnosperms

are generally more drought resistant as a result of lower stomatal sensitivity to vapor pressure deficit (VPD) and more cavitation-resistant xylem (Carnicer et al., 2013). Conifer xylem is made up entirely of tracheids, whereas angiosperms may produce both tracheids and wide vessels, which have higher hydraulic conductivity, but a smaller margin of safety with regard to xylem pressures (Carnicer et al., 2013). In addition, angiosperms have more complex anatomical responses to drought, such as changes in vessel connectivity (Brodribb et al., 2012). Moreover, much research attention has been given to drought tolerance in conifers because they are a dominant component of many arid zone forests. We first address the definitions of 'drought tolerance' and the physiological mechanisms involved. We then describe three major methods that have been used to study drought tolerance genetics, and review the major findings to date. Finally, we discuss how genetic tools may aid forest management, and needs for future research. In particular, we recommend the combination of complementary methods, and the broadening of the range of phenotypes, taxa, life stages and timespans examined.

II. Drought tolerance

1. Definition of 'drought tolerance'

A basic definition of drought tolerance is the ability to survive, and sometimes grow, during periods of water shortage. Survival and growth are often correlated, with trees exhibiting a history of below average growth or abrupt decreases in growth having higher mortality (Wyckoff & Clark, 2002; van Mantgem et al., 2003; Das et al., 2007; Huang et al., 2015). Because of this, and because tree genetics studies are often motivated by wood production, some studies define drought tolerance as growth maintenance (e.g. Orwig & Abrams, 1997; Eilmann et al., 2010). However, drought length and duration can affect the growth-mortality relationship. For instance, growth plasticity (high growth in good conditions, low in bad) may be adaptive in variable environments (Santos-Del-Blanco et al., 2013). Populations adapted to extended drought often exhibit conservative resource use strategies, resulting in slower growth rates, even in favorable conditions (Correia et al., 2008; Herrero et al., 2013; Kerr et al., 2015; Montwé et al., 2015).

Drought tolerance can be broken down into several categories. 'Drought avoidance' strategies (e.g. deep roots or stomatal adjustments) reduce exposure to drought stress (McDowell *et al.*, 2008). However, it is unclear to what degree trees truly avoid drought stress relative to plants that go dormant. 'Drought resistance' is the ability to withstand drought exposure, whereas 'drought resilience' is a measure of how quickly a tree can resume normal growth when conditions improve (Lloret *et al.*, 2011; Eilmann & Rigling, 2012).

2. Mechanisms of drought tolerance

The physiological responses of trees to drought have been reviewed elsewhere (Sala *et al.*, 2010; McDowell, 2011; Anderegg *et al.*, 2012; Choat *et al.*, 2012; Carnicer *et al.*, 2013; Sevanto *et al.*, 2014). We include a brief discussion here to highlight some of the traits and processes that are involved in drought avoidance, resistance and resilience.

Avoidance of hydraulic failure and decreased cellular water potential Conifers manage tissue water potential in two main ways: isohydric trees close stomata to maintain water potential, whereas anisohydric species allow water potential to drop (Tardieu & Simonneau, 1998) (Fig. 1a,b). Isohydric trees use increasing abscisic acid (ABA) concentrations as a signal to keep stomata closed, whereas anisohydric trees use low leaf water potential itself as a signal to close stomata (Brodribb *et al.*, 2014). Anisohydric conifers include many Cupressaceae and some Taxaceae (McDowell *et al.*, 2008; Brodribb *et al.*, 2014).

Xylem architecture affects how changes in stomatal conductivity influence cavitation risk, and anisohydric trees tend to have xylem that is more cavitation resistant (McDowell *et al.*, 2008). Wider tracheids increase conductivity and the risk of hydraulic failure (Sperry *et al.*, 2006), whereas those with smaller inter-tracheid pits or more lignified walls are less vulnerable (Sperry *et al.*, 1988; Tyree & Ewers, 1991). The reduction of leaf area with branch die-back, reduced needle number or smaller needles (Cinnirella *et al.*, 2002; Eldhuset *et al.*, 2013) can also reduce water loss. Anisohydric species often exhibit branch die-back during drought, whereas isohydric trees typically retain a full canopy until death (Koepke *et al.*, 2010) (Fig. 1a,b).

Some conifer species can refill xylem following cavitation. This is thought to be an energy-intensive process that depends on carbon reserves (Carnicer *et al.*, 2013). This may explain why drought-stressed trees can exhibit lower refilling capability (Cinnirella *et al.*, 2002). *Picea abies* refills freezing-cavitated xylem before soils have thawed by taking up water through its needles (Mayr *et al.*, 2014). This could explain why other conifers can refill xylem in the absence of positive root pressures, unlike co-occurring angiosperms (Sperry *et al.*, 1994). However, refilled xylem may be less resistant to future drought stress, a characteristic known as 'cavitation fatigue' (Hacke *et al.*, 2001).

Prevention of cellular damage when water content drops Loss of water potential in cells is associated with cell turgor loss, denaturation of proteins and changes in membrane fluidity. To avoid cellular damage, plants synthesize molecules that act as osmotic balancing agents. These reduce cellular solute potential, and may increase turgor at lower water potentials. In addition, hydrophilic compounds can prevent the membranes from leaking (Valliyodan & Nguyen, 2006; Yuxiu et al., 2007). Other compounds stabilize proteins or detoxify reactive oxygen species. These protective molecules include proteins such as chaperonins and dehydrins (Hamanishi & Campbell, 2011), the amino acid proline (Hayat et al., 2012) and various carbohydrates (Valliyodan & Nguyen, 2006; Lorenz et al., 2011). We hypothesize that protective molecules may be produced earlier during a drought in anisohydric species because leaf water potential drops more quickly (Fig. 1c).

Management of carbon reserves As a result of reduced stomatal conductance (g_s), oxidative damage and other factors, photosynthetic rates and chlorophyll concentrations often decline during drought (Dubos *et al.*, 2003; Watkinson *et al.*, 2003). Therefore, in addition to protecting cells from damage, increased allocation to

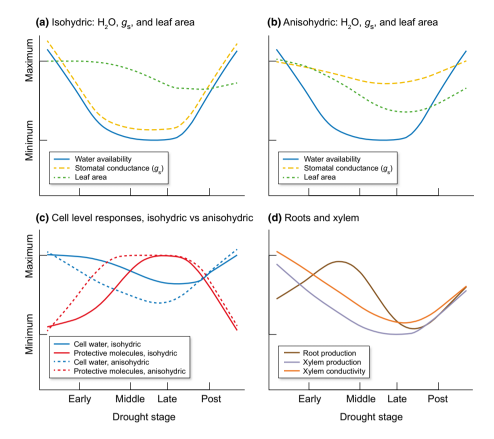


Fig. 1 Hypothetical drought responses. (a) In isohydric trees, stomatal conductance is reduced when water availability is low to avoid water loss, but leaf area is largely maintained. (b) In anisohydric trees, stomatal conductance is maintained, but leaf area may be reduced. (c) As a result, cell water content drops more strongly and earlier in anisohydric species. This could necessitate earlier expression of protective molecules. (d) Trees sometimes produce more roots in response to early or mild drought, but prolonged drought can reduce root growth. Xylem production and the conductivity of the xylem tend to decline during drought.

nonstructural carbohydrates (NSCs) may help to avoid carbon starvation by keeping energy resources in easily mobilizable forms. The patterns of change in overall NSCs and starches seem to differ between species and drought length and severity (Oberhuber *et al.*, 2011; Adams *et al.*, 2013; Anderegg & Anderegg, 2013; Reinhardt *et al.*, 2015). Clear evidence of death as a result of carbon depletion is still lacking (McDowell & Sevanto, 2010; Sala *et al.*, 2010). However, carbon storage and allocation patterns do vary under drought stress (Galiano Pérez *et al.*, 2017), demonstrating implications of water limitation on carbon availability.

Influences of anatomical structures, individual plasticity and drought legacy effects Root and branch patterns can influence the degree of water stress experienced. The number of branches and leaves affects total transpiration. Rooting depth affects access to deep soil water and is probably crucial for seedlings (Cregg & Zhang, 2001; Olmo et al., 2014) as well as adult trees in areas with seasonal drought (O'Brien et al., 2017). Deep roots may also redistribute water from deep to shallow soils (Horton & Hart, 1998; Brooks et al., 2002). More small diameter roots, with high surface area: volume and a lower vulnerability to cavitation, may aid drought resistance (Phillips et al., 2016).

Structural changes can have long-lasting effects. Decreasing soil moisture can induce greater root production, but extended drought reduces root mass (Eldhuset *et al.*, 2013) (Fig. 1d), which limits responsiveness to precipitation pulses (Plaut *et al.*, 2013). Lumen width and cell wall thickness of tracheids are plastic, with those produced in moist seasons and years generally being wider, more numerous and thinner walled than those

produced in dry periods (Eldhuset *et al.*, 2013; Xu *et al.*, 2014) (Fig. 1d). Xylem is often functional for multiple years (Eilmann *et al.*, 2010; Mayr *et al.*, 2014; Xu *et al.*, 2014; Kimak & Leuenberger, 2015), and so current drought responses can affect water transport during future drought.

The production of protective molecules typically drops soon after normal water potential is restored (Lorenz *et al.*, 2011). However, transcriptional and physiological 'memory' in stomatal guard cells has been observed, with stressed plants maintaining smaller stomatal apertures when re-watered (Virlouvet & Fromm, 2015). There may also be 'legacy effects' on NSC production (Villar-Salvador *et al.*, 1999) and traits such as growth and xylem anatomy (Anderegg *et al.*, 2015). Plants that quickly return to normal could gain a growth advantage. In areas in which recurring drought is common, however, we hypothesize that this memory effect reduces mortality risk.

There are multiple traits involved at different stages of the drought response (Fig. 2). Stomatal control and patterns of root and shoot growth affect the degree to which a plant avoids drought stress. These traits plus xylem morphology, protective molecule production, changes in carbohydrate metabolism and pathogen defenses influence drought resistance. Finally, the recovery rate of photosynthesis and other processes, the degree of persistent changes in structure and the ability to refill xylem affect drought resilience. In the next two sections, we first review the methods used to date to examine genetic controls on ecologically important traits, and then explore how these methods have been and can be leveraged to test for genetic variation in, and identify the genetic basis of, the traits and processes addressed above.

Physiological studies		Genetic studies					
Key traits/ responses	Stages	Expression	Provenance	Genome scan/ G2E	QTL/G2P		
Stomatal control (ABA pathway)	Avoidance Resistance Resilience	1	1?	Δ	Δ		
Shoot growth	Avoidance Resistance Resilience	? (cell division)	1?	Δ ? (cell division)	Δ ? (cell division)		
Protective molecules (e.g. dehydrins)	Resistance	1	?	Δ	Δ		
Pathogen defense	Resistance	1	?	Δ	?		
Nonstructural carbohydrates	Resistance Resilience	Δ ? (carbohydrate metabolism)	?	Δ ? (carbohydrate metabolism)	?		
Root growth	Avoidance Resistance Resilience	?	† ?	?	?		
Xylem morphology	Resistance Resilience	?	?	Δ ? (cell wall)	Δ ? (cell wall)		
Xylem refilling	Resilience	?	?	?	?		

Fig. 2 Connections between traits or responses identified in physiological studies as important for drought tolerance and genetic studies. The latter include shifts in gene expression under drought, differences between source populations (provenances) from dry vs wet environments, differences in alleles across environmental gradients (genome scan or G2E association), and differences in alleles associated with a phenotype (QTL or G2P association). Arrows indicate the direction of difference between dry vs wet, environment or provenance; Δ indicate a difference where the direction of change is unclear. Question marks indicate where a relationship is unclear, either because it has not been investigated, or because it is unclear how or if the genes identified relate to the process or trait. Narrow arrows paired with question marks indicate a directional change in something that might be related to the process or trait.

III. Investigation of the genetic basis of drought tolerance: three main approaches

1. Gene expression/transcriptome studies

Gene expression or transcriptome studies examine changes in the amount of RNA transcripts to identify genes that are upregulated or downregulated under different conditions. Changes in the amount of a gene product (e.g. chaperone protein) can result in different phenotypic responses, even if all individuals have the same gene sequence. Such changes are responsible for plasticity, and may involve temporary or heritable epigenetic modifications (Bräutigam *et al.*, 2013).

Gene expression studies may involve a variety of techniques, but most recent studies have used microarray chips – DNA probes to which cDNA or RNA hybridize, resulting in fluorescence (Watkinson et al., 2003; Lorenz et al., 2011) - or cDNA sequencing (e.g. RNAseq) (Lorenz et al., 2006; Behringer et al., 2015). The latter avoids the need for probe and microarray design and can survey whole novel transcriptomes (Wang et al., 2009). Real-time quantitative polymerase chain reaction (qPCR) is highly sensitive, but is most often used to target specific candidate genes (Perdiguero et al., 2012a; Velasco-Conde et al., 2012) or to confirm a subset of expression changes (Perdiguero et al., 2012b; Behringer et al., 2015). All techniques are sensitive to which tissues are sampled at what time (Fossdal et al., 2007; Yeaman et al., 2014). Moreover, unless expression responses in different genotypes or populations are explicitly compared, this approach does not address local adaptation.

2. Provenance studies

Provenance or common garden studies, where seedlings from many different sources are planted in a common environment, began to reveal heritable differences between tree populations long before the availability of genetic marker data (Callaham & Liddicoet, 1961; Rehfeldt et al., 1984; Rehfeldt, 1991; Correia et al., 2008). Provenance studies established in the mid-20th century to identify seed zones for replanting or highly productive genotypes have been re-purposed to investigate potential responses to climate change (Schmidtling, 1994; Rehfeldt et al., 2002, 2014; Wang et al., 2006; O'Neill et al., 2008; Leites et al., 2012). Many recent studies have also used seedling common gardens (see Section IV.2). Studies conducted across multiple sites, or incorporating multiple treatments, can estimate the plasticity of traits, allowing the fitting of transfer functions that predict performance based on source and planting environments (Wang et al., 2006; Leites et al., 2012; Rehfeldt et al., 2014). However, such studies do not reveal which genes are responsible for observed differences unless paired with other techniques. It should be noted that there is usually substantial variation within tree populations (Aitken et al., 2008). The third set of approaches can be used to investigate the causes of heritable variation between populations and individuals.

3. Quantitative trait locus (QTL), genome scan and genotype association studies

These approaches aim to identify genes or genomic regions related to a trait or to adaptation along environmental gradients. QTL

studies are a classic way to identify the loci involved in continuous trait variation. However, although QTLs for a number of traits have been identified in trees, this approach has had limited success for a variety of reasons, many of which are reviewed in González-Martínez *et al.* (2006). For instance, a great deal of time and space is needed to cross parental tree lines and raise a sufficient sample size of progeny. Conifers also have very large genomes with low linkage disequilibrium (LD) and, without enough genetic markers available, most QTLs are undetectable (Neale & Kremer, 2011). In addition, high-resolution genetic/physical maps or positional cloning is needed to identify causal genes/mutations (Aitken *et al.*, 2008; Neale & Kremer, 2011; de Miguel *et al.*, 2014).

By contrast, genome scan and association studies make use of large numbers of newly available markers (e.g. single nucleotide polymorphisms, SNPs), and are carried out in highly diverse outcrossing natural populations (González-Martínez *et al.*, 2006). Genome scans identify loci that differ more or less between populations than expected by chance (Eveno *et al.*, 2008; Namroud *et al.*, 2008; Prunier *et al.*, 2011). For instance, outlier F_{st} values can be used to infer the type of selection: balancing selection results in low F_{st} and shared alleles, and divergent selection in high F_{st} with segregated alleles. Genome scans can also identify patterns suggestive of a selective sweep. These studies do not automatically provide information about which, if any, environmental variables are responsible for the pattern. One can test whether patterns of differentiation match an environmental gradient, but this is necessarily a *post-hoc* interpretation (Eckert *et al.*, 2010a).

Association studies use a regression approach to identify loci in which genetic variation is associated with variation in trait values or home environment. Such analyses can be carried out at the individual (genotype) or population (allele frequency) level. Genotype-to-environment (G2E) association studies identify loci that vary along environmental gradients (Eckert *et al.*, 2010a; Prunier *et al.*, 2011; Frichot *et al.*, 2013). An association between an SNP and aridity, for example, suggests that the gene or its regulatory region affects performance in wet vs dry environments. This does not reveal *how* the locus affects phenotype, and careful interpretation is needed as a result of correlation between climatic variables. Genotype-to-phenotype (G2P) association studies identify loci correlated with a particular phenotype (Neale & Savolainen, 2004; Holliday *et al.*, 2010; Cumbie *et al.*, 2011), but the phenotype may or may not be relevant for fitness in the field.

Most association studies in conifers to date have used SNPs in a limited number of candidate genes (Gonzalez-Martinez et al., 2005; Aitken et al., 2008; Holliday et al., 2010). This ensures that genes suspected of involvement are surveyed, but limits the ability to identify additional loci. However, with the decreasing cost of sequencing, approaches that generate large numbers of SNPs are increasingly being used for genome-wide association studies (GWAS). One set of approaches, including RAD-seq and genotyping-by-sequencing (GBS), involves the use of restriction enzymes to cut and sequence a small subset of the genome (Elshire et al., 2011; Poland & Rife, 2012; Andrews et al., 2016). This can produce tens of thousands of SNPs with high coverage (Chen et al., 2013; Karam et al., 2015; Pan et al., 2015). Many of these SNPs will be in noncoding regions, which is good for the potential

discovery of regulatory regions, but can limit the number of gene associations detected. Another approach involves the creation of a transcriptome or full genome sequence for a species, and the development of probes for all or most of the putative genes to identify SNPs (Cokus *et al.*, 2015; Jaramillo-Correa *et al.*, 2015; Gugger *et al.*, 2016). This approach can also yield useful gene expression data if multiple tissue types or treatments are included in the development of the transcriptome (Yeaman *et al.*, 2014).

IV. Conifer drought tolerance genetics: what we know

1. Changes in gene expression during drought

Overall changes in gene expression in drought-stressed conifer seedlings Most drought gene expression studies in conifers have focused on pine seedlings, with a few investigating other Pinaceae genera (Table 1, column 3). The direction of expression responses to the environment, including dry conditions, is highly conserved (74% of orthologs) between *Pinus contorta* and *Picea glauca* × *engelmannii*, even though average expression levels often differ (Yeaman *et al.*, 2014). It is unclear whether this is true across conifer families. No expression studies have focused on adult drought responses.

The methods used to induce drought stress vary. Studies have withheld water for a specified period (Perdiguero *et al.*, 2012a), until soil moisture reached a threshold (Velasco-Conde *et al.*, 2012; Eldhuset *et al.*, 2013) or needles wilted (Lorenz *et al.*, 2011), or needle water content declined to a certain level (Watkinson *et al.*, 2003; Behringer *et al.*, 2015). Some have used chemically induced water stress (Dubos *et al.*, 2003; Perdiguero *et al.*, 2012b). Caution must therefore be used in interpreting differences across studies, as these could be methodological artifacts (Watkinson *et al.*, 2003).

Genes related to signaling and gene transcription are frequently upregulated in drought-stressed seedlings. Changes in signal cascades must precede changes in their targets, and such expression shifts often occur within the first week of drought stress. Those in the ABA pathway are well represented (Fig. 2; Table 1). In addition to being involved in stomatal closure, ABA signaling can affect shoot growth and water uptake (Parent *et al.*, 2009; Hamanishi & Campbell, 2011). However, there are also ABA-independent pathways in most taxa, which may use leaf water potential as a signal (Valliyodan & Nguyen, 2006; Hamanishi & Campbell, 2011; Brodribb *et al.*, 2014). Upregulation of genes in the ethylene pathway (Lorenz *et al.*, 2011; Perdiguero *et al.*, 2012b) could be related to reduced shoot growth or leaf area (Carnicer *et al.*, 2013).

Genes related to protective molecules are also frequently upregulated (Fig. 2; Table 1). Late-embryogenesis-abundant (LEA) proteins, named for their role in seeds, appear to stabilize proteins and membranes and prevent protein aggregation (Close, 1996; Goyal *et al.*, 2005). Dehydrins, a subgroup of LEAs, often protect against drought stress, although some are induced by other abiotic stresses (Yuxiu *et al.*, 2007; Perdiguero *et al.*, 2012a; Velasco-Conde *et al.*, 2012). Heat shock proteins, detoxification enzymes and genes in the synthesis and transport pathways of osmoprotective carbohydrates and proline may also be upregulated

Table 1 Gene categories linked to drought responses in conifers by four different approaches

Functional category	Subcategory	Expression studies	Genome scans	G2E association	QTL or G2P association
Protective proteins	Dehydrins	ABAL ¹ ; PIPI ^{14,17} ; PITA ^{11,12,18}	PIPI ⁸		PITA ⁹
L	Other LEAs	ABAL ¹ ; PIPI ¹⁵ ; PITA ^{11,12,18} (+/-)	PIMA ¹⁶		
	Heat shock proteins	PIPI ¹⁵ ; PITA ^{12,18}		PICE & PIMU ¹³	PIPI ²
	Chaperones & their regulators	,		PITA ⁵	PIPI ²
Osmoprotective	Transport	PIPI ¹⁵ ; PITA ¹²			PILA ⁷
carbohydrates	Synthases	ABAL ¹ ; PIPI ¹⁵ ; PITA ¹²			
Proline synthesis	.,	PITA ¹²			
Detoxification	Aldehyde dehydrogenase	PIPI ¹⁵		PIPI ¹⁰	
	Oxidative stress defense	PIPI ¹⁵ ; PITA ¹²		PITA ⁵	PIPI ² ; PITA ⁹
	Other	ABAL ¹ ; PITA ¹⁸			,
Signaling & gene	ABA pathway	ABAL ¹ (+/-); PITA ^{11,12}	PIPI ⁸	PICE ¹³	PITA ³
transcription	Ethylene	PIPI ¹⁵ ; PITA ¹²		PIPI ¹⁰	PIPI ²
	Various transcription factors	ABAL ¹ (+/-); PIPI ¹⁵ ; PITA ¹²	PIMA ¹⁶	PIPI ¹⁰	PIPI ² ; PITA ⁹
	Kinases	ABAL ¹ ; PIPI ^{4,15} (–)		PILA ⁷ ; PIMU ¹³	PIPI ²
		· · · · · · · · · · · · · · · · · · ·		PITA ⁵	
	Non-kinase signal tranduction	ABAL ¹ ; PIPI ¹⁵		PICE ¹³	PIPI ²
	Histones	PITA ¹²			
	Methylation	PITA ¹¹	PIPI ⁸		
	Other	PIPI ^{4,15} (+/-); PITA ¹¹		PILA ⁷ ; PITA ⁵	PIPI ² ; PITA ³
Cell wall construction	Other	ABAL ¹ (-); PIPI ⁴ (-); PITA ^{11,12} (-)	PIPI ⁸	LADE & PIMU ¹³ ; PIPI ¹⁰	PIPI ² ; PITA ⁹
Cytoskeleton		ABAL ¹ (–); PITA ¹⁸		PIPI ¹⁰ ; PITA ⁵	1111,1117
Growth/cell division		PIPI ^{4,15} (–); PITA ¹²		PITA ⁵	PIPI ²
or expansion		1111 (-),111A		1117	1 11 1
Cation/H ⁺ transporters		ABAL ¹		PIPI ¹⁰ ; PITA ⁵	PIPI ²
Carbohydrate	Carbohydrate synthesis	ABAL ¹ (-); $PIPI^{4,15}$ (+/-)		1111 ,1117	1 11 1
metabolism	Breakdown	ABAL ¹ ; PIPI ¹⁵ ; PITA ^{12,18} (+/-)		LADE & PIMU ¹³	PIPI ²
metabolism	Other	PIPI ⁴ (+/-); PITA ¹² (-)		LADE & PIMU ¹³	1 11 1
	Other	1111 (17-),11174 (-)		PILA ⁷	
Biotic stress defense		ABAL ¹ ; PIPI ^{4,15} ; PITA ^{11,12}		PILA ⁷ ; PITA ⁵	
Protein handling/	Ubiquitin pathway	ABAL ¹	PIMA ¹⁵	PIMU ¹³ ; PITA ^{5,6}	PIPI ²
breakdown	Lysosome	PIPI ⁴ (–)	PIIVIA	LADE ¹³	PIPI
Dreakdown	Proteasome	PIPI (-) PIPI ¹⁵		PICE ¹³	
Toomalakian		PIPI		LADE & PICE ¹³ ; PIPI ¹⁰	
Translation	Ribosomes or mRNA processing tRNAs			ABAL ¹³	
		PIPI ⁴		ADAL	
Disates with a sign	Translation initiation	PIPI PIPI ⁴ (—)		PIPI ¹⁰ ; PITA ⁵	PIPI ²
Photosynthesis	Photosystems	PIPI (-) PIPI ⁴ (+/-)		PIPI ; PITA	PIPI
	Photorespiration	PIPI (+/-) PIPI ⁴ (-); PITA ¹⁸			
	Rubisco	PIPI (-); PITA			PIPI ²
District and the literature	Other chloroplast proteins	ADAL 1. DIDI15. DITA12 ()			PIPI ²
Lipid metabolism	6	ABAL ¹ ; PIPI ¹⁵ ; PITA ¹² (+/-)		PICE ¹³ ; PIPI ¹⁰ PITA ⁵	PIPI~
Transport	Carbohydrate	ABAL ¹		PICE'S; PIPI'S PITAS	DIDI?
	Water (aquaporins)	ABAL ¹ ; PITA ¹²			PIPI ²
	Protein	PIPI ⁴ (-)			
	Other	PIPI ¹⁵			DIDI ²
Amino acid metabolism		ABAL ¹ ; PIPI ¹⁵ ; PITA ¹⁸ (+/-)		DIDI ¹⁰ DIT 45	PIPI ²
Mitochondrial proteins		PITA ¹²		PIPI ¹⁰ ; PITA ⁵	PIPI ² ; PITA ³
Heavy metal binding		PIPI ⁴ (–)			
Pollen				PITA ⁵	
Miscellaneous metabolism		ABAL ¹ (+/-); PITA ¹¹		PITA ⁵	

^(–) In expression studies indicates downregulation as opposed to upregulation, whereas (+/–) indicates a difference between tissues or genes in the category. ABA, abscisic acid; G2E, genotype-to-environment; G2P, genotype-to-phenotype; LEA, late embryogenesis abundant; QTL, quantitative trait locus; WUE, intrinsic water use efficiency. Species codes: ABAL, Abies alba; LADE, Larix decidua; PIAB, Picea abies; PICE, Pinus cembra; PILA, Pinus lambertiana; PIMA, Picea mariana; PIMU, Pinus mugo; PIPI, Pinus pinaster; PITA, Pinus taeda.

¹Behringer *et al.* (2015) (qPCR-validated loci with 3–10-fold change in expression); ²de Miguel *et al.* (2014) (QTLs for traits measured during drought stress); ³Cumbie *et al.* (2011); ⁴Dubos *et al.* (2003); ⁵Eckert *et al.* (2010a) (loci with Bayes Factor > 100); ⁶Eckert *et al.* (2010b); ⁷Eckert *et al.* (2015); ⁸Eveno *et al.* (2008); ⁹Gonzalez-Martinez *et al.* (2008); ¹⁰Jaramillo-Correa *et al.* (2015); ¹¹Lorenz *et al.* (2006); ¹²Lorenz *et al.* (2011) (genes with greatest change in expression); ¹³Mosca *et al.* (2012); ¹⁴Perdiguero *et al.* (2012a); ¹⁵Perdiguero *et al.* (2012b); ¹⁶Prunier *et al.* (2011); ¹⁷Velasco-Conde *et al.* (2012); ¹⁸Watkinson *et al.* (2003).

(Lorenz et al., 2011; Perdiguero et al., 2012b; Behringer et al., 2015).

Genes involved in pathogen or biotic stress defenses are often upregulated during drought stress, but those involved in growth, including cell division and wall construction, are often downregulated (Dubos *et al.*, 2003; Lorenz *et al.*, 2006, 2011; Perdiguero *et al.*, 2012b; Behringer *et al.*, 2015) (Fig. 2; Table 1). Up or down changes in carbohydrate and lipid metabolism and protein handling pathways are also evident, although these are more difficult to interpret. Aquaporins, which affect membrane water permeability, were found to be upregulated in two studies (Lorenz *et al.*, 2011; Behringer *et al.*, 2015).

When drought-stressed seedlings are re-watered, most gene expression quickly returns to normal. In *Pinus taeda*, only 76 of the 2445 genes with altered expression during drought were still different after 48 h of recovery (Lorenz *et al.*, 2011). Lorenz *et al.* (2006) found 11 genes upregulated in 'recovered' *P. taeda* seedlings relative to either drought-stressed or well-watered seedlings, including probable cell wall proteins, an aquaporin and a gene involved in vacuole function. These may reflect recovery or repair processes that occur once drought stress is removed.

Studies targeting specific gene families or functions Gene families illustrate the complexity of expression responses to drought. *Pinus pinaster* has at least eight dehydrin genes, based on expressed sequence tag (EST) analyses (Perdiguero *et al.*, 2012a). Three of five were downregulated during drought, whereas the other two were upregulated (Velasco-Conde *et al.*, 2012). Most dehydrin induction occurred after 20 d of drought (Perdiguero *et al.*, 2012a), which may be why a similar but shorter study (Perdiguero *et al.*, 2012b) did not reveal the upregulation of dehydrins. Expression can also vary by tissue. Of seven dehydrins examined in *P. abies*, drought stress upregulated four in needles, but only two in bark, with one being downregulated in bark (Eldhuset *et al.*, 2013).

To investigate the link between drought and defense gene expression, Fossdal *et al.* (2007) exposed *P. abies* seedlings to a pathogen (*Rhizoctonia*), drought stress or both, and examined the transcription of 14 candidate defense genes. Genes were upregulated more slowly in drought-stressed seedlings than in pathogen-inoculated seedlings. The combined treatment led to more rapid and/or higher expression of many defense genes than either alone, which may be related to the synergistic mortality risks posed by biotic and abiotic stressors. Pleiotropic effects for some drought/ defense-related genes are also possible, but none have been identified to date.

2. Local adaptation to drought in conifers

Provenance studies Multiple provenance studies have identified patterns consistent with local adaptation to drought. Trees from drier climates often exhibit conservative growth strategies (Fig. 2), such as slower height or needle growth (de la Mata *et al.*, 2014), less aboveground biomass or a shorter growing season (Kerr *et al.*, 2015). Seedlings from dry environments often also exhibit more root growth (Fig. 2) and higher drought survival (Cregg & Zhang,

2001; Matías et al., 2014; Kolb et al., 2016). Provenance trials of *Pinus halepensis* have shown mixed responses, with low growth and high water use efficiency (WUE) in dry-sourced populations (Voltas et al., 2008), but high growth in populations from intermediate-aridity areas (Klein et al., 2013), which may be related to growth plasticity.

Because of the importance of carbon resources for plants, WUE - the ratio of carbon fixed to water lost - has long been considered to be closely tied to drought tolerance (Farquhar et al., 1989; Cregg & Zhang, 2001). Measures that integrate over longer time periods, such as the carbon isotope ratio δ^{13} C (Farguhar *et al.*, 1989; Livingston et al., 1999), are most frequently used to represent changes in WUE in trees. However, although different measures of WUE are often correlated (de Miguel et al., 2014), they are not interchangeable. For example, carbon discrimination (Δ) is sensitive to chloroplast carbon concentrations and mesophyll conductance, whereas WUE itself is heavily influenced by evaporative demand, which does not directly affect Δ (Seibt et al., 2008). In addition, nitrogen fertilization can increase WUE and decrease Δ , but does not affect g_s or transpiration (Ripullone *et al.*, 2004). Thus, WUE and Δ do not always co-vary, and caution is needed in the interpretation of δ^{13} C as a measure of WUE. Additional caution is warranted when using WUE as an indication of drought tolerance. High WUE may not be adaptive in some dry environments if the use of less water per unit carbon fixed does not result in slower depletion of soil water (as a result of competition or other factors), or if plants with higher WUE grow faster and thus use more total water.

Although a few studies have shown higher δ^{13} C for populations from dry sites (Kerr *et al.*, 2015), others have shown the opposite (Guy & Holowachuk, 2001; Lamy *et al.*, 2011). There was no difference between three populations of *Pinus ponderosa* seedlings from varying climates in δ^{13} C or instantaneous WUE; the drought-adapted populations exhibited greater plasticity in water use (Zhang *et al.*, 1997). In *P. halepensis*, however, individuals from more mesic sources showed higher plasticity of WUE than those from drier sources (Klein *et al.*, 2013), but dry sources may show higher average WUE (Voltas *et al.*, 2008). Highly plastic growth and water usage reduce apparent WUE over the whole season compared with consistently moderate to low water usage (Zhang *et al.*, 1997; Kerr *et al.*, 2015).

Instantaneous measures of WUE can change over a day, whereas integrated measures can differ significantly for a source population grown under different conditions or for the same tree across years (Zhang et al., 1997; Klein et al., 2013). Changes in WUE may thus be a useful indication of drought stress, but, in conifers, radial growth and WUE are often weakly or negatively correlated (Adams & Kolb, 2004; Correia et al., 2008; Eilmann et al., 2010; Marguerit et al., 2014). In pines, higher WUE usually results from reduced gs (Fig. 2) and/or reduced leaf area (Zhang et al., 1997; de Miguel et al., 2012, 2014; Marguerit et al., 2014; Reinhardt et al., 2015), which can limit photosynthesis and growth (Brendel et al., 2002). Low gs can also result in higher tissue temperatures, which can be damaging, particularly in seedlings (Kolb & Robberecht, 1996).

Drought length and severity can influence measures of relative drought tolerance between populations. In *P. ponderosa* seedlings,

the relative growth rate under moist conditions was positively correlated with previously measured tolerance to severe drought, whereas, under 4-wk drought, the intermediate-drought-tolerant population grew faster (Zhang et al., 1997). When Silim et al. (2001) examined Picea sitchensis, P. glauca and their hybrids, they found that P. sitchensis and the hybrids had the highest WUE and growth in well-watered conditions, but P. glauca and the hybrids had higher WUE and growth in drought conditions. Similarly, the relative transpiration and photosynthetic rates, WUE and growth of P. halepensis tree provenances differed between near-desert and Mediterranean planting sites (Atzmon et al., 2004). Such shifts in ranking are often a result of plasticity differences between populations.

Provenances of *P. pinaster* from across the species' range did not vary in cavitation resistance, suggesting uniform selection or lack of genetic variation (Lamy *et al.*, 2011). In *P. halepensis*, however, the percentage loss of conductivity (attributed to cavitation) differed significantly between provenances, but not between environments (Klein *et al.*, 2013). Although plasticity has been observed in xylem wall thickening, time to thickening and number of cells in *Picea mariana* in drought experiments (Balducci *et al.*, 2013), cell anatomy studies often focus on only one population, so that the extent of local adaptation is unknown.

Genetic signals of local adaptation Genome scans have identified loci in conifers that may be under differential selection across environments (Fig. 2; Table 1). Of 13 candidate genes for drought response in P. pinaster, two showed signs of divergent selection, although only one (a cell wall gene) exhibited a pattern correlated with a climatic gradient; three, including two dehydrins, showed evidence of balancing selection (Eveno et al., 2008). Prunier et al. (2011) examined SNPs from 313 candidate genes in P. mariana and found 16 that exhibited differentiation correlated with precipitation, including a LEA protein and genes in the ubiquitin protein handling pathway. However, differentiation between populations can be driven by processes unrelated to climatic gradients. Conifer populations are likely to violate the assumptions of such tests because they rarely form discrete isolated populations and are often far from demographic equilibrium; mis-specification of population hierarchical structure can lead to high false positive rates (Eckert et al., 2010b). However, newer methods are being developed that avoid frequently violated assumptions and reduce false positives (Whitlock & Lotterhos, 2015).

The approach preferred by recent studies is to directly assess the association of loci with environmental gradients (G2E), whilst controlling for population structure (Fig. 2; Table 1). Jaramillo-Correa et al. (2015) examined the correlation of P. pinaster candidate gene SNP allele frequencies with climate principal component (PC) axes, using transcriptome-wide SNPs to control for population structure and demographic history. They identified 18 environmentally associated SNPs, many of which were in genes relating to carbohydrate transport, cell wall construction and photosynthesis. Two surveys of P. taeda examined associations between candidate gene SNPs and environmental gradients. One examined the association of these loci with five climatic PC axes

(Eckert *et al.*, 2010a), whereas the other used an aridity index (precipitation/PET) for each county (Eckert *et al.*, 2010b). There was some overlap in function between the loci identified (Table 1). However, the studies disagreed on whether SNPs associated with climate also tended to be $F_{\rm st}$ outliers.

G2E associations have been detected even over short geographical distances, suggesting that selection can drive local adaptation in the presence of high gene flow. Eckert *et al.* (2015) examined *Pinus lambertiana* populations around Lake Tahoe, an area of c. $35 \times 65 \, \mathrm{km}^2$, and found 11 genes associated with environmental PCs reflecting differences in water availability. These included genes involved in carbohydrate metabolism and transport and response to biotic stress (Table 1).

A cross-species comparison of environmental associations suggests some similarities in the genetic mechanisms involved in climatic tolerances across conifer genera. For each of four European conifer species in the Italian Alps, 6–18 SNPs (of 249–693 investigated) were associated with precipitation/temperature PC axes (Table 1). There was some overlap between species in the genes represented, including heat shock proteins, and cell wall construction and carbohydrate metabolism genes (Mosca *et al.*, 2012).

3. Linking genes to traits and traits to drought adaptation by combining approaches

Gene expression studies have identified a range of genes that may be involved in drought responses, but these results are not easily connected to the results of physiological or provenance response studies. First, RNA transcripts reflect the genes being expressed at a particular instant, whereas morphological or physiological traits are the result of processes acting over a longer time. Second, most gene expression studies do not examine differences between populations. Although some evidence suggests that stronger gene expression changes during stress are associated with greater growth or survival, different genotypes and demographic stages can show significant differences in gene expression changes (Hamanishi & Campbell, 2011).

A few studies have begun to address this. Provenances of *P. pinaster* differed in the expression response of two dehydrin genes, as well as in physiology and mortality rates (Velasco-Conde *et al.*, 2012). Similarly, three genotypes of *P. taeda* differed in their gene expression responses to drought and re-watering (Lorenz *et al.*, 2006). More such studies are needed, but care must be taken to distinguish between drivers of expression differences. For instance, a more drought-sensitive tree might express higher levels of dehydrins at a given drought stage because the leaf water potential has dropped faster than in a drought-resistant tree, whereas the resistant tree might express higher levels of dehydrins than the sensitive tree at a given leaf water potential.

Genome scan and G2E association studies can be useful tools in the search for genes responsible for local adaptation. Although such studies can identify loci at which allele frequencies differ between environments, it is not always clear how these differences are connected to phenotypic differences, and thus what traits are under selection in a given environment. This is where QTL and G2P association studies are useful.

Most conifer QTL studies have focused on wood traits, growth or yield. Of the two that have examined drought tolerance, the first identified four significant and four suggestive QTLs for δ^{13} C in *P. pinaster*, none of which co-located with QTLs for ring width (Brendel *et al.*, 2002). The second examined a wider range of traits – photosynthesis (A_n), chlorophyll fluorescence, g_s , δ^{13} C, intrinsic WUE (WUE_i) and specific leaf area (SLA) – in F₁ cross seedlings of *P. pinaster* when well watered or after 1 or 2 wk without water, and identified 28 significant and 27 suggestive QTLs (de Miguel *et al.*, 2014). Locations of the QTLs for each trait (except SLA) varied by time point. Candidate genes within the QTLs were identified (Table 1): those for g_s and WUE_i included stomatal regulation, ABA signaling and cell wall construction genes; those for δ^{13} C included an aquaporin; and those for chlorophyll fluorescence included transcription factors and a histone chaperone.

G2P studies focusing on quantitative traits (including cold hardiness, budset date, yield and wood properties) have generally been successful in identifying associated loci (Holliday et al., 2010; Resende et al., 2012; Gamal El-Dien et al., 2015). However, only a few studies have investigated drought tolerance in conifers (Fig. 2; Table 1), with less success. All such studies used δ^{13} C as the focal trait. As we argue in Section VI, other traits would probably yield results that are more helpful for the understanding of drought responses. Gonzalez-Martinez et al. (2008) examined 41 candidate stress response genes of P. taeda, using 61 tree families planted at two sites. However, drought stress was probably mild, and they only identified one strongly associated gene and one weakly associated gene at each site. A later study on the same species examining 3938 SNPs identified seven new associations with δ^{13} C (Cumbic *et al.*, 2011). Four of the associations were with unknown proteins, with only a transcription factor probably involved in the ABA-mediated stress response having an obvious connection to drought responses.

G2P and G2E association studies complement one another, with the first identifying loci linked to targeted traits, but not whether these loci are under selection in nature, and the second doing the opposite. The combination of these approaches is useful for the identification of genes and traits under selection in natural settings, but so far few studies have taken this approach. Eckert et al. (2015) tested the association of SNPs with five phenotypic traits and 11 environmental variables across 10 P. lambertiana populations around Lake Tahoe. This study identified six genes associated with phenotypic traits (bud flush date, needle nitrogen, δ^{13} C and height growth), and 31 associated with environmental PCs. Two genes were associated with both a trait and an environmental axis, including a glucose transport protein associated with δ^{13} C and environmental variables linked to water availability. A study focusing on multiple drought response traits and a larger number of SNPs might be able to identify more genes that have variants associated with both environmental gradients and drought tolerance traits.

Some traits and processes involved in drought response have been better studied at the genetic level than others (Fig. 2). Provenance studies have indicated that differences in stomatal control and shoot growth (physiologically related to all three drought response stages) are often involved in local adaptation to drought, and all other study types have identified the genes likely to be involved (related to the ABA pathway and cell division, respectively). However, although root growth has also been identified as important by provenance studies, root-growth-related genes have not been identified. Conversely, although genes related to resistance traits, such as changes in carbohydrate metabolism, and protective and pathogen defense molecule production, have been identified in expression or association studies, these traits have been largely ignored in provenance studies. Finally, xylem traits, including refilling ability, have not been the focus of any genetic study type.

V. Implications for the management of forests in a changing world

Tree improvement programs that aim to increase growth potential and stress resistance face the challenges of long generation times, the need for large-scale field experiments and the late expression of traits such as wood density (Isik, 2014; Gamal El-Dien *et al.*, 2015). Genomic selection, already routinely used in livestock breeding, has been proposed as a method of speeding up this process by using marker-predicted breeding values for phenotypes of interest (Grattapaglia & Resende, 2011; Isik, 2014). This approach is suitable for species with low LD and for traits with complex genetic architectures as it uses thousands of markers with effects that are estimated simultaneously (Gamal El-Dien *et al.*, 2015). As with traditional phenotypic selection, accuracy is likely to be greatest when tests are carried out in environments similar to the target environment, because of the high likelihood of genotype × environment interactions (Isik, 2014).

Several recent studies have demonstrated the potential of genomic selection approaches for traits of interest to forestry. Resende *et al.* (2012) carried out an early evaluation of genomic selection in *P. taeda*, making use of clonally replicated individuals grown on four sites and genotyped at 4825 SNPs. They found that the accuracy of prediction models within sites ranged from 0.63 to 0.75 for diameter and height, and estimated that the breeding cycle could be speeded up by 50% with this method. Gamal El-Dien *et al.* (2015) used GBS to genotype over 1000 interior spruce trees (*P. glauca* × *P. engelmanii*) over three sites that had been phenotyped for yield and wood attributes, and found that the incorporation of genomic information produced more accurate heritability estimates. Genomic estimated breeding values were most accurate (0.47–0.77) when data from multiple sites were used to fit the model.

Of even more relevance to selection for drought tolerance, Jaramillo-Correa *et al.* (2015) identified 18 SNPs associated with climatic PC axes in *P. pinaster*, and found that the frequency of locally advantageous alleles at these loci correlated with population-level survival rates in a common garden at the hot/dry end of the species range. Together with the growth trait analyses, these results suggest that association techniques could be applied to predict breeding values for overall drought tolerance or particular drought tolerance traits even though only some of the loci involved have been identified.

There is evidence of significant potential for selection approaches to improve drought responses in conifers. Provenance studies have

shown evidence of genetic differentiation between populations in drought responses, and genome scan and G2E associations are finding evidence of natural selection on within-species genetic variation. Second, heritabilities for drought tolerance traits, when these have been examined, tend to be moderate to high. The calculation of heritability requires pedigree information: parentoffspring or sibling and half-sib comparisons. Narrow-sense heritability is the fraction of the variance in a trait attributable to additive genetic variation, as opposed to environmental and nonadditive genetic variation. Because heritability depends on both genetic variation in the population assessed and the degree of variation caused by the environment, estimates are not transferable between situations. In *P. pinaster*, estimates of δ^{13} C narrow-sense heritability ranged from 0.17 to 0.41, depending on how many individuals of what populations were assessed in what sites; and ring width (0.19-0.25) and height growth rates (0.35) were also moderately heritable (Brendel et al., 2002; Lamy et al., 2011; Marguerit et al., 2014). In the same species, heritability of P₅₀ (a measure of cavitation resistance) was 0.44, but this was driven more strongly by low levels of other sources of variation rather than high additive genetic variation (Lamy et al., 2011). Across species, measured heritabilities for δ^{13} C range from the very high 0.7 for *Araucaria cunninghamii* to < 0.1 for *P. taeda* (Johnsen *et al.*, 1999; Prasolova et al., 2001; Baltunis et al., 2008).

Managers of wild forests are often focused on ensuring the resilience and function of the ecosystem rather than productivity. G2E and G2P association studies may help to identify seed sources that could be 'preadapted' to projected conditions for replanting in wildlands. However, wild trees face a range of challenges, including disease and competition, as well as drought (Sthultz et al., 2009; Grady et al., 2015). Stand structure (McDowell et al., 2006; Das et al., 2008, 2011; Fernandez et al., 2012) and soil properties (Koepke et al., 2010; Phillips et al., 2016) may also directly affect how trees experience drought stress. Studies that integrate standlevel processes with genetic testing can further bridge gaps between genetic experiments and forest-scale management. Restoration projects could be used as experiments (Howe & Martínez-Garza, 2014) to test genomic predictions of survival and growth in a given environment, as well as the effects of genetic composition and diversity of the planted population on restoration success.

VI. Future directions

In order to address the remaining questions relating to the genetics of drought tolerance in conifers, we offer several recommendations for future studies.

1. More 'crossover' between the major categories of drought genetics studies

Common garden, gene expression and genetic association studies all have different strengths and weaknesses, and none alone will answer the question of how genetic differences affect drought tolerance (Fig. 3). As described previously, a combination of different types of association study may help to identify loci that are under selection in the wild and the traits they influence. Similarly,

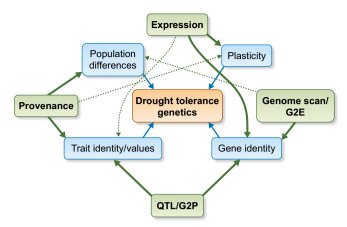


Fig. 3 The study types used to investigate drought tolerance genetics to date (green boxes) each address only some of the components (blue boxes) needed to understand the links between genes, phenotypes and fitness in a given environment. Some components are always addressed (solid line arrows), whereas others are sometimes addressed (dashed line arrows). A combination of study types can allow a more complete set of connections to be made: for instance, a combination of provenance and expression approaches can be used to investigate the role of expression differences in local adaptation. G2E, genotype-to-environment; G2P, genotype-to-phenotype; QTL, quantitative trait locus.

gene expression studies could easily be combined with common garden studies of adults or seedlings to address whether differences in drought responses between populations or genotypes are a result of differences in gene sequences, gene expression patterns or both.

2. Less focus on WUE

Many studies to date have focused on WUE, often using δ^{13} C as a proxy. As discussed above, however, WUE is a ratio of changes in photosynthesis and transpiration, which can both vary, and higher WUE may or may not be associated with greater survival or growth in dry conditions. Moreover, different measures of WUE are not entirely consistent. We therefore recommend that future studies use survival and/or growth during and following drought as the metric of overall 'drought tolerance', and measure photosynthesis and water loss separately if these are processes of interest. The time involved in the measurement of traits for hundreds or thousands of individuals has encouraged the focus on easily measured δ^{13} C, but much progress has been made in high-throughput phenotyping techniques (Plomion et al., 2016). For instance, thermal and longwave infrared sensors can measure leaf temperature or stomatal conductance, near and short-wave infrared sensors can measure leaf water content, and fluorescence sensors can measure chlorophyll content and photosystem efficiency (Araus & Cairns, 2014; Fahlgren et al., 2015).

3. More focus on carbohydrate metabolism, xylem refilling and root growth

There are several traits and processes that have been suggested to be important for drought response by physiological studies, but about which there is little genetic information (Fig. 2). Genetic studies

frequently identify genes related to carbohydrate metabolism and transport as having altered expression or allele frequencies depending on water availability. It is difficult to make sense of these patterns because the link between these metabolic changes and tree function and survival during drought is still unclear. We also know relatively little about which species can refill cavitated xylem, under what circumstances and by what mechanisms. Thus, it is difficult to determine whether any genes identified by expression or G2E studies are involved in this process. Similarly, how roots and root growth respond to changes in water availability, and what genes are involved in these responses, remain poorly understood. Although the measurement of root architecture can be complex, high-throughput methods are being developed for this as well (Araus & Cairns, 2014).

4. More experiments using adult trees and longer timespans

Most experimental studies, including those looking at gene expression, have focused on seedlings. There may be important differences in how different life stages respond to drought. For instance, Pinus nigra ssp. laricio adults have been observed to follow an isohydric strategy, whereas seedlings in a glasshouse experiment did not (Cinnirella et al., 2002). Although it is more complicated to impose drought treatments on adults, drought experiments have been carried out on adult trees using networks of rain shields/ gutters to intercept precipitation and direct it away from the trees (Borghetti et al., 1998; Cinnirella et al., 2002; Plaut et al., 2013). This water can also be re-directed to other plots to create 'wellwatered' treatments. For the most part, these studies have been carried out on natural populations. However, if they were coupled with provenance study plantings, one could test for population or genotypic differences in adult drought response. Likewise, apart from some long-term provenance studies (Atzmon et al., 2004),

most experiments span a few days to a few months. In order to investigate drought resilience and legacy effects, more multi-year studies are needed.

5. Explicit consideration of different types of drought in the context of natural environments

The length and intensity of drought can affect which trait combinations result in greater fitness. In Section IV.1, we mentioned the great diversity of methods used to induce or measure drought stress treatments in gene expression studies. The same diversity is found in G2P and provenance studies as well. There is a need to assess: (1) whether environmental treatments (e.g. watering frequency or soil moisture) roughly match the range of conditions in the environments in which the target species does or might grow; (2) how environmental treatments relate to plant stress measures (e.g. leaf water content or wilting); and (3) whether traits, responses or genotypes associated with drought tolerance in the glasshouse or laboratory predict performance in the field. In addition, studies testing longer term drought treatments are lacking, as are those that explicitly test variable combinations of drought length and severity. Future work should address these gaps.

6. Identification of the role of 'mystery' genes

In most of the genetic studies cited above, a relatively high proportion of the genes expressed or linked to phenotypes or environmental gradients of interest either have unknown or poorly defined functions. Behringer *et al.* (2015), for instance, found that, of the 832 transcripts analyzed for gene ontology, 538 either had no database hits or could not be assigned to a biological process. Although this could be partly addressed with further studies in

Table 2 Frequency of inclusion of conifer taxa in drought tolerance and genomic studies, as well as features that influence their attractiveness for such studies

Conifer taxon	Representation in drought tolerance studies	Genomic information ¹	Aridity of natural habitat	Economic importance
Pinaceae				
Pinus	High, particularly Pinus pinaster	Full genome: <i>Pinus taeda</i> , <i>Pinus lambertiana</i> . Partial info for many	Mesic to xeric	Generally fast-growing. Plantations worldwide for pulp and wood
Picea	Moderate	Full genome: Pinus abies and Pinus glauca	Boreal areas (cold water limitation)	Cool climate plantations for pulp and wood
Pseudotsuga	Low	Full genome for <i>Pinus</i> menziesii	Mesic	Plantations for pulp and wood in Europe and North America
Other (e.g. <i>Larix</i> , Abies, Tsuga, Cedrus)	Extremely low	Larix siberica sequence in progress	Variable – <i>Cedrus</i> xeric, <i>Tsuga</i> mesic, etc.	Some high – <i>Larix</i> often planted, <i>Cedrus</i> wood historically valued
Cupressaceae	Low	Some candidate gene work in redwood, candidate genes and linkage maps in Cryptomeria japonica	Many mesic, but some Cuppressoideae (Calocedrus, Cupressus, Juniperus) inhabit xeric areas and many species anisohydric	Slow-growing. Only a few (e.g. redwood, Sequoia sempervirens) plantation grown for timber
Other families (e.g. Araucariaceae, Podocarpaceae)	Extremely low	Little	Variable, but often mesic (some tropical)	Some high-value timbers, esp. Araucariacea. Infrequently planted for wood or pulp

¹http://www.pinegenome.org/projects.php

traditional model organisms, such as *Arabidopsis*, analysis of loblolly pine and Norway and Sitka spruce genome sequences suggests that there could be thousands of conifer-specific gene families (De La Torre *et al.*, 2014). This shortcoming must be addressed by further development of model systems in conifers.

7. Inclusion of overlooked conifer taxa

Unsurprisingly, the conifer taxa that have received the most attention in terms of drought tolerance studies or genomic studies are those that are of high economic value, especially those that are frequently grown in plantations. Species from xeric environments also tend to be over-represented in drought tolerance studies. This means that most drought tolerance genetics studies have been carried out on pines (Pinus), with a modest representation of spruce (Picea) and Douglas-fir (Pseudotsuga) (Table 2). Although this focus is understandable, this means that a number of ecologically or economically significant taxa have been left out, notably the Cupressaceae (Brodribb et al., 2014). Many Juniperus and New World Cupressus and Calocedrus species are impressively drought tolerant. In the well-studied pinyon-juniper woodlands of the American Southwest, anisohydric juniper tends to exhibit lower mortality than Pinus edulis during severe drought (McDowell et al., 2008; Koepke et al., 2010). However, not all pine species exhibit isohydric behavior (Cinnirella et al., 2002). Thus, the genes involved in drought response could differ substantially between families, genera or species.

VII. Conclusion

Conifer responses to drought can involve a variety of morphological and physiological traits. Provenance studies, gene expression studies and genomic approaches all have different strengths and weaknesses, and have revealed different aspects of how conifers respond and adapt to drought stress. The genes and pathways that are commonly involved include those related to ABA signaling, carbohydrate metabolism, the production of proteins and other molecules that protect cells from changes in osmotic potential or oxidative stress, and defense against biotic threats that might take advantage of a drought-stressed plant. In some cases, the same genes appear to be involved in the responses of distantly related species. However, not all traits thought to be involved in drought tolerance have been investigated from a genetic perspective, and drought response strategies are likely to differ between conifer species, populations and life stages in ways that have not been fully explored. The impact of these responses on fitness in turn depends on the intensity and duration of the drought stress. Further progress will require the combination of insights from these disparate approaches. Comparisons across species, families and life stages will probably yield insights into which responses are most adaptive under different circumstances. Nevertheless, early attempts at predicting phenotypes from genotypes suggest that genetic tools may be able to aid managers to select appropriate planting stock in the near future, at least for the better studied species.

References

- Adams HD, Germino MJ, Breshears DD, Barron-Gafford GA, Guardiola-Claramonte M, Zou CB, Huxman TE. 2013. Nonstructural leaf carbohydrate dynamics of *Pinus edulis* during drought-induced tree mortality reveal role for carbon metabolism in mortality mechanism. *New Phytologist* 197: 1142–1151.
- Adams HD, Kolb TE. 2004. Drought responses of conifers in ecotone forests of northern Arizona: tree ring growth and leaf δ^{13} C. *Oecologia* 140: 217–225.
- Aitken SN, Yeaman S, Holliday JA, Wang T, Curtis-McLane S. 2008. Adaptation, migration, or extirpation: climate change outcomes for tree populations. *Evolutionary Applications* 1: 95–111.
- Allen CD, Macalady AK, Chenchouni H, Bachelet D, McDowell N, Vennetier M, Kitzberger T, Rigling A, Breshears DD, Hogg EH (Ted) et al. 2010. A global overview of drought and heat-induced tree mortality reveals emerging climate change risks for forests. Forest Ecology and Management 259: 660–684.
- Anderegg WRL, Anderegg LDL. 2013. Hydraulic and carbohydrate changes in experimental drought-induced mortality of saplings in two conifer species. *Tree Physiology* 33: 252–260.
- Anderegg WRL, Berry JA, Field CB. 2012. Linking definitions, mechanisms, and modeling of drought-induced tree death. *Trends in Plant Science* 17: 693–700.
- Anderegg WRL, Schwalm C, Biondi F, Camarero JJ, Koch G, Litvak M, Ogle K, Shaw JD, Shevliakova E, Williams AP et al. 2015. Pervasive drought legacies in forest ecosystems and their implications for carbon cycle models. Science 349: 528–532.
- Andrews KR, Good JM, Miller MR, Luikart G, Hohenlohe PA. 2016. Harnessing the power of RADseq for ecological and evolutionary genomics. *Nature Reviews Genetics* 17: 81–92.
- Araus JL, Cairns JE. 2014. Field high-throughput phenotyping: the new crop breeding frontier. *Trends in Plant Science* 19: 52–61.
- Atzmon N, Moshe Y, Schiller G. 2004. Ecophysiological response to severe drought in *Pinus halepensis* Mill. trees of two provenances. *Plant Ecology* 171: 15–22.
- Balducci L, Deslauriers A, Giovannelli A, Rossi S, Rathgeber CBK. 2013. Effects of temperature and water deficit on cambial activity and woody ring features in *Picea mariana* saplings. *Tree Physiology* 33: 1006–1017.
- Baltunis BS, Martin TA, Huber DA, Davis JM. 2008. Inheritance of foliar stable carbon isotope discrimination and third-year height in *Pinus taeda* clones on contrasting sites in Florida and Georgia. *Tree Genetics & Genomes* 4: 797–807
- Behringer D, Zimmermann H, Ziegenhagen B, Liepelt S. 2015. Differential gene expression reveals candidate genes for drought stress response in *Abies alba* (Pinaceae). *PLoS ONE* 10: e0124564.
- Borghetti M, Cinnirella S, Magnani F, Saracino A. 1998. Impact of long-term drought on xylem embolism and growth in *Pinus halepensis* Mill. *Trees* 12: 187–195.
- Bräutigam K, Vining KJ, Lafon-Placette C, Fossdal CG, Mirouze M, Marcos JG, Fluch S, Fraga MF, Guevara MÁ, Abarca D et al. 2013. Epigenetic regulation of adaptive responses of forest tree species to the environment. Ecology and Evolution 3: 399–415.
- Brendel O, Pot D, Plomion C, Rozenberg P, Guehl J-M. 2002. Genetic parameters and QTL analysis of δ^{13} C and ring width in maritime pine. *Plant, Cell & Environment* 25: 945–953.
- Breshears DD, Cobb NS, Rich PM, Price KP, Allen CD, Balice RG, Romme WH, Kastens JH, Floyd ML, Belnap J et al. 2005. Regional vegetation die-off in response to global-change-type drought. Proceedings of the National Academy of Sciences, USA 102: 15144–15148.
- Brodribb TJ, McAdam SAM, Jordan GJ, Martins SCV. 2014. Conifer species adapt to low-rainfall climates by following one of two divergent pathways. *Proceedings of the National Academy of Sciences, USA* 111: 14489–14493.
- Brodribb TJ, Pittermann J, Coomes DA. 2012. Elegance versus speed: examining the competition between conifer and angiosperm trees. *International Journal of Plant Sciences* 173: 673–694.
- Brooks JR, Meinzer FC, Coulombe R, Gregg J. 2002. Hydraulic redistribution of soil water during summer drought in two contrasting Pacific Northwest coniferous forests. *Tree Physiology* 22: 1107–1117.
- Callaham RZ, Liddicoet AR. 1961. Altitudinal variation at 20 years in Ponderosa and Jeffrey pines. *Journal of Forestry* 59: 814–820.

- Carnicer J, Barbeta A, Sperlich D, Coll M, Peñuelas J. 2013. Contrasting trait syndromes in angiosperms and conifers are associated with different responses of tree growth to temperature on a large scale. *Frontiers in Plant Science* 4: 1–19.
- Chen C, Mitchell SE, Elshire RJ, Buckler ES, El-Kassaby YA. 2013. Mining conifers' mega-genome using rapid and efficient multiplexed high-throughput genotyping-by-sequencing (GBS) SNP discovery platform. *Tree Genetics & Genomes* 9: 1537–1544.
- Choat B, Jansen S, Brodribb TJ, Cochard H, Delzon S, Bhaskar R, Bucci SJ, Feild TS, Gleason SM, Hacke UG et al. 2012. Global convergence in the vulnerability of forests to drought. Nature 491: 752–755.
- Cinnirella S, Magnani F, Saracino A, Borghetti M. 2002. Response of a mature *Pinus laricio* plantation to a three-year restriction of water supply: structural and functional acclimation to drought. *Tree Physiology* 22: 21–30.
- Close TJ. 1996. Dehydrins: emergence of a biochemical role of a family of plant dehydration proteins. *Physiologia Plantarum* 97: 795–803.
- Cokus SJ, Gugger PF, Sork VL. 2015. Evolutionary insights from *de novo* transcriptome assembly and SNP discovery in California white oaks. *BMC Genomics* 16: 552–569.
- Correia I, Almeida MH, Aguiar A, Alía R, David TS, Pereira JS. 2008. Variations in growth, survival and carbon isotope composition (δ¹³C) among *Pinus pinaster* populations of different geographic origins. *Tree Physiology* 28: 1545–1552.
- Cregg BM, Zhang JW. 2001. Physiology and morphology of *Pinus sylvestris* seedlings from diverse sources under cyclic drought stress. *Forest Ecology and Management* 154: 131–139.
- Cumbie WP, Eckert A, Wegrzyn J, Whetten R, Neale D, Goldfarb B. 2011.

 Association genetics of carbon isotope discrimination, height and foliar nitrogen in a natural population of *Pinus taeda* L. *Heredity* 107: 105–114.
- Das A, Battles J, Stephenson NL, van Mantgem PJ. 2011. The contribution of competition to tree mortality in old-growth coniferous forests. Forest Ecology and Management 261: 1203–1213.
- Das A, Battles J, van Mantgem PJ, Stephenson NL. 2008. Spatial elements of mortality risk in old-growth forests. *Ecology* 89: 1744–1756.
- Das AJ, Battles JJ, Stephenson NL, van Mantgem PJ. 2007. The relationship between tree growth patterns and likelihood of mortality: a study of two tree species in the Sierra Nevada. Canadian Journal of Forest Research 37: 580–597.
- De La Torre AR, Birol I, Bousquet J, Ingvarsson PK, Jansson S, Jones SJM, Keeling CI, MacKay J, Nilsson O, Ritland K *et al.* 2014. Insights into conifer gigagenomes. *Plant Physiology* 166: 1724–1732.
- Dubos C, Le Provost G, Pot D, Salin F, Lalane C, Madur D, Frigerio J-M, Plomion C. 2003. Identification and characterization of water-stress-responsive genes in hydroponically grown maritime pine (*Pinus pinaster*) seedlings. *Tree Physiology* 23: 169–179.
- Eckert AJ, Bower AD, González-Martínez SC, Wegrzyn JL, Coop G, Neale DB. **2010a**. Back to nature: ecological genomics of loblolly pine (*Pinus taeda*, Pinaceae). *Molecular Ecology* **19**: 3789–3805.
- Eckert AJ, van Heerwaarden J, Wegrzyn JL, Nelson CD, Ross-Ibarra J, Gonzalez-Martinez SC, Neale DB. 2010b. Patterns of population structure and environmental associations to aridity across the range of loblolly pine (*Pinus taeda* L., Pinaceae). *Genetics* 185: 969–982.
- Eckert AJ, Maloney PE, Vogler DR, Jensen CE, Mix AD, Neale DB. 2015. Local adaptation at fine spatial scales: an example from sugar pine (*Pinus lambertiana*, Pinaceae). *Tree Genetics & Genomes* 11: 42–58.
- Eilmann B, Buchmann N, Siegwolf R, Saurer M, Cherubini P, Rigling A. 2010. Fast response of Scots pine to improved water availability reflected in tree-ring width and δ^{13} C: fast response of Scots pine to improved water availability. *Plant, Cell & Environment* 33: 1351–1360.
- Eilmann B, Rigling A. 2012. Tree-growth analyses to estimate tree species' drought tolerance. *Tree Physiology* 32: 178–187.
- Eldhuset TD, Nagy NE, Volařík D, Børja I, Gebauer R, Yakovlev IA, Krokene P. 2013. Drought affects tracheid structure, dehydrin expression, and above- and belowground growth in 5-year-old Norway spruce. *Plant and Soil* 366: 305–320.
- Elshire RJ, Glaubitz JC, Sun Q, Poland JA, Kawamoto K, Buckler ES, Mitchell SE. 2011. A robust, simple genotyping-by-sequencing (GBS) approach for high diversity species. PLoS ONE 6: e19379.
- Eveno E, Collada C, Guevara MA, Leger V, Soto A, Diaz L, Leger P, Gonzalez-Martinez SC, Cervera MT, Plomion C et al. 2008. Contrasting patterns of

- selection at *Pinus pinaster* Ait. drought stress candidate genes as revealed by genetic differentiation analyses. *Molecular Biology and Evolution* **25**: 417–437.
- Fahlgren N, Gehan MA, Baxter I. 2015. Lights, camera, action: high-throughput plant phenotyping is ready for a close-up. *Current Opinion in Plant Biology* 24: 93–99.
- Farquhar GD, Ehleringer JR, Hubick KT. 1989. Carbon isotope discrimination and photosynthesis. *Annual Review of Plant Physiology and Plant Molecular Biology* 40: 503–537
- Fernandez M, Gyenge J, de Urquiza M, Varela S. 2012. Adaptability to climate change in forestry species: drought effects on growth and wood anatomy of ponderosa pines growing at different competition levels. *Forest Systems* 21: 162–173
- Fossdal CG, Nagy NE, Johnsen Ø, Dalen LS. 2007. Local and systemic stress responses in Norway spruce: similarities in gene expression between a compatible pathogen interaction and drought stress. *Physiological and Molecular Plant Pathology* 70: 161–173.
- Frichot E, Schoville SD, Bouchard G, François O. 2013. Testing for associations between loci and environmental gradients using latent factor mixed models. *Molecular Biology and Evolution* 30: 1687–1699.
- Gailing O, Vornam B, Leinemann L, Finkeldey R. 2009. Genetic and genomic approaches to assess adaptive genetic variation in plants: forest trees as a model. *Physiologia Plantarum* 137: 509–519.
- Galiano Pérez L, Timofeeva G, Saurer M, Siegwolf R, Martínez-Vilalta J, Hommel R, Gessler A. 2017. The fate of recently fixed carbon after drought release: towards unravelling C storage regulation in *Tilia platyphyllos* and *Pinus sylvestris. Plant, Cell & Environment* 40: 1711–1724.
- Gamal El-Dien O, Ratcliffe B, Klápště J, Chen C, Porth I, El-Kassaby YA. 2015. Prediction accuracies for growth and wood attributes of interior spruce in space using genotyping-by-sequencing. *BMC Genomics* 16: 370–385.
- Gonzalez-Martinez SC, Ersoz E, Brown G, Wheeler NC, Neale DB. 2005. DNA sequence variation and selection of tag single-nucleotide polymorphisms at candidate genes for drought-stress response in *Pinus taeda* L. *Genetics* 172: 1915–1926.
- Gonzalez-Martinez SC, Huber D, Ersoz E, Davis JM, Neale DB. 2008. Association genetics in *Pinus taeda* L. II. Carbon isotope discrimination. *Heredity* 101: 19–26.
- González-Martínez SC, Krutovsky KV, Neale DB. 2006. Forest-tree population genomics and adaptive evolution. *New Phytologist* 170: 227–238.
- Goyal K, Walton LJ, Tunnacliffe A. 2005. LEA proteins prevent protein aggregation due to water stress. The Biochemical Journal 388: 151–157.
- Grady KC, Kolb TE, Ikeda DH, Whitham TG. 2015. A bridge too far: cold and pathogen constraints to assisted migration of riparian forests. *Restoration Ecology* 23: 811–820.
- Grattapaglia D, Resende MDV. 2011. Genomic selection in forest tree breeding. Tree Genetics & Genomes 7: 241–255.
- Gugger PF, Cokus SJ, Sork VL. 2016. Association of transcriptome-wide sequence variation with climate gradients in valley oak (*Quercus lobata*). *Tree Genetics & Genomes* 12: 15–28.
- Guy RD, Holowachuk DL. 2001. Population differences in stable carbon isotope ratio of *Pinus contorta* Dougl. ex Loud.: relationship to environment, climate of origin, and growth potential. *Canadian Journal of Botany* 79: 274–283.
- Hacke UG, Stiller V, Sperry JS, Pittermann J, McCulloh KA. 2001. Cavitation fatigue. Embolism and refilling cycles can weaken the cavitation resistance of xylem. *Plant Physiology* 125: 779–786.
- Hamanishi ET, Campbell MM. 2011. Genome-wide responses to drought in forest trees. Forestry 84: 273–283.
- Hayat S, Hayat Q, Alyemeni MN, Wani AS, Pichtel J, Ahmad A. 2012. Role of proline under changing environments. *Plant Signaling & Behavior* 7: 1456–1466.
- Herrero A, Castro J, Zamora R, Delgado-Huertas A, Querejeta JI. 2013. Growth and stable isotope signals associated with drought-related mortality in saplings of two coexisting pine species. *Oecologia* 173: 1613–1624.
- Holliday JA, Ritland K, Aitken SN. 2010. Widespread, ecologically relevant genetic markers developed from association mapping of climate-related traits in Sitka spruce (*Picea sitchensis*). New Phytologist 188: 501–514.
- Horton JL, Hart SC. 1998. Hydraulic lift: a potentially important ecosystem process. Trends in Ecology & Evolution 13: 232–235.
- Howe H, Martínez-Garza C. 2014. Restoration as experiment. *Botanical Sciences* 92: 459–468.

- Huang M, Piao S, Sun Y, Ciais P, Cheng L, Mao J, Poulter B, Shi X, Zeng Z, Wang Y. 2015. Change in terrestrial ecosystem water-use efficiency over the last three decades. Global Change Biology 21: 2366–2378.
- IPCC. 2013. Stocker TF, Qin D, Plattner G-K, Tignor M, Allen SK, Boschung J, Nauels A, Xia Y, Bex V, Midgley PM, eds. Climate change 2013: the physical science basis. Contribution of Working Group I to the fifth assessment report of the Intergovernmental Panel on Climate Change. Cambridge, UK & New York, NY, USA: Cambridge University Press.
- Isik F. 2014. Genomic selection in forest tree breeding: the concept and an outlook to the future. New Forests 45: 379–401.
- Jaramillo-Correa J-P, Rodriguez-Quilon I, Grivet D, Lepoittevin C, Sebastiani F, Heuertz M, Garnier-Gere PH, Alia R, Plomion C, Vendramin GG et al. 2015. Molecular proxies for climate maladaptation in a long-lived tree (*Pinus pinaster* Aiton, Pinaceae). Genetics 199: 793–807.
- Johnsen KH, Flanagan LB, Huber DA, Major JE. 1999. Genetic variation in growth, carbon isotope discrimination, and foliar N concentration in *Picea mariana*: analyses from a half-diallel mating design using field-grown trees. Canadian Journal of Forest Research 29: 1727–1735.
- Karam M-J, Lefèvre F, Dagher-Kharrat MB, Pinosio S, Vendramin G. 2015.
 Genomic exploration and molecular marker development in a large and complex conifer genome using RADseq and mRNAseq. *Molecular Ecology Resources* 15: 601–612.
- Kerr KL, Meinzer FC, McCulloh KA, Woodruff DR, Marias DE. 2015. Expression of functional traits during seedling establishment in two populations of *Pinus ponderosa* from contrasting climates. *Tree Physiology* 35: 535–548.
- Kimak A, Leuenberger M. 2015. Are carbohydrate storage strategies of trees traceable by early—latewood carbon isotope differences? Trees 29: 859–870.
- Klein T, Di Matteo G, Rotenberg E, Cohen S, Yakir D. 2013. Differential ecophysiological response of a major Mediterranean pine species across a climatic gradient. *Tree Physiology* 33: 26–36.
- Koepke DF, Kolb TE, Adams HD. 2010. Variation in woody plant mortality and dieback from severe drought among soils, plant groups, and species within a northern Arizona ecotone. *Oecologia* 163: 1079–1090.
- Kolb PF, Robberecht R. 1996. High temperature and drought stress effects on survival of *Pinus ponderosa* seedlings. *Tree Physiology* 16: 665–672.
- Kolb TE, Grady KC, McEttrick MP, Herrero A. 2016. Local-scale drought adaptation of ponderosa pine seedlings at habitat ecotones. Forest Science 62: 641– 651.
- Lamy J-B, Bouffier L, Burlett R, Plomion C, Cochard H, Delzon S. 2011. Uniform selection as a primary force reducing population genetic differentiation of cavitation resistance across a species range. PLoS ONE 6: e23476.
- Leites LP, Rehfeldt GE, Robinson AP, Crookston NL, Jaquish B. 2012.
 Possibilities and limitations of using historic provenance tests to infer forest species growth responses to climate change. *Natural Resource Modeling* 25: 409–433.
- Livingston NJ, Guy RD, Sun ZJ, Ethier GJ. 1999. The effects of nitrogen stress on the stable carbon isotope composition, productivity and water use efficiency of white spruce (*Picea glauca* (Moench) Voss) seedlings. *Plant, Cell & Environment* 22: 281–289.
- Lloret F, Keeling EG, Sala A. 2011. Components of tree resilience: effects of successive low-growth episodes in old ponderosa pine forests. Oikos 120: 1909– 1920
- Lorenz WW, Alba R, Yu Y-S, Bordeaux JM, Simões M, Dean JF. 2011. Microarray analysis and scale-free gene networks identify candidate regulators in droughtstressed roots of loblolly pine (*P. taeda* L.). *BMC Genomics* 12: 264–280.
- Lorenz WW, Sun F, Liang C, Kolychev D, Wang H, Zhao X, Cordonnier-Pratt M-M, Pratt LH, Dean JF. 2006. Water stress-responsive genes in loblolly pine (*Pinus taeda*) roots identified by analyses of expressed sequence tag libraries. *Tree Physiology* 26: 1–16.
- van Mantgem PJ, Stephenson NL. 2007. Apparent climatically induced increase of tree mortality rates in a temperate forest. Ecology Letters 10: 909–916.
- van Mantgem PJ, Stephenson NL, Mutch LS, Johnson VG, Esperanza AM, Parsons DJ. 2003. Growth rate predicts mortality of *Abies concolor* in both burned and unburned stands. *Canadian Journal of Forest Research* 33: 1029–1038.
- Marguerit E, Bouffier L, Chancerel E, Costa P, Lagane F, Guehl J-M, Plomion C, Brendel O. 2014. The genetics of water-use efficiency and its relation to growth in maritime pine. *Journal of Experimental Botany* 65: 4757–4768.

- de la Mata R, Merlo E, Zas R. 2014. Among-population variation and plasticity to drought of Atlantic, Mediterranean, and interprovenance hybrid populations of maritime pine. *Tree Genetics & Genomes* 10: 1191–1203.
- Matías L, González-Díaz P, Jump AS. 2014. Larger investment in roots in southern range-edge populations of Scots pine is associated with increased growth and seedling resistance to extreme drought in response to simulated climate change. *Environmental and Experimental Botany* 105: 32–38.
- Mayr S, Schmid P, Laur J, Rosner S, Charra-Vaskou K, Dämon B, Hacke UG. 2014. Uptake of water via branches helps timberline conifers refill embolized xylem in late winter. *Plant Physiology* 164: 1731–1740.
- McDowell N, Allen CD, Marshall L. 2010. Growth, carbon-isotope discrimination, and drought-associated mortality across a *Pinus ponderosa* elevational transect. *Global Change Biology* 16: 399–415.
- McDowell N, Pockman WT, Allen CD, Breshears DD, Cobb N, Kolb T, Plaut J, Sperry J, West A, Williams DG *et al.* 2008. Mechanisms of plant survival and mortality during drought: why do some plants survive while others succumb to drought? *New Phytologist* 178: 719–739.
- McDowell N, Sevanto S. 2010. The mechanisms of carbon starvation: how, when, or does it even occur at all? *New Phytologist* 186: 263–264.
- McDowell NG. 2011. Mechanisms linking drought, hydraulics, carbon metabolism, and vegetation mortality. *Plant Physiology* 155: 1051–1059.
- McDowell NG, Adams HD, Bailey JD, Hess M, Kolb TE. 2006. Homeostatic maintenance of ponderosa pine gas exchange in response to stand density changes. *Ecological Applications* 16: 1164–1182.
- de Miguel M, Cabezas J-A, de María N, Sánchez-Gómez D, Guevara MÁ, Vélez MD, Sáez-Laguna E, Díaz LM, Mancha JA, Barbero MC et al. 2014. Genetic control of functional traits related to photosynthesis and water use efficiency in Pinus pinaster Ait. drought response: integration of genome annotation, allele association and QTL detection for candidate gene identification. BMC Genomics 15: 464.
- de Miguel M, Sanchez-Gomez D, Cervera MT, Aranda I. 2012. Functional and genetic characterization of gas exchange and intrinsic water use efficiency in a fullsib family of *Pinus pinaster* Ait. in response to drought. *Tree Physiology* 32: 94– 103
- Montwé D, Spiecker H, Hamann A. 2015. Five decades of growth in a genetic field trial of Douglas-fir reveal trade-offs between productivity and drought tolerance. *Tree Genetics & Genomes* 11: 29–39.
- Mosca E, Eckert AJ, Di Pierro EA, Rocchini D, La Porta N, Belletti P, Neale DB. 2012. The geographical and environmental determinants of genetic diversity for four alpine conifers of the European Alps. *Molecular Ecology* 21: 5530–5545.
- Namroud M-C, Beaulieu J, Juge N, Laroche J, Bousquet J. 2008. Scanning the genome for gene single nucleotide polymorphisms involved in adaptive population differentiation in white spruce. *Molecular Ecology* 17: 3599–3613.
- Neale DB, Kremer A. 2011. Forest tree genomics: growing resources and applications. *Nature Reviews Genetics* 12: 111–122.
- Neale DB, Savolainen O. 2004. Association genetics of complex traits in conifers. Trends in Plant Science 9: 325–330.
- Oberhuber W, Swidrak I, Pirkebner D, Gruber A. 2011. Temporal dynamics of nonstructural carbohydrates and xylem growth in *Pinus sylvestris* exposed to drought. *Canadian Journal of Forest Research* 41: 1590–1597.
- O'Brien MJ, Engelbrecht BMJ, Joswig J, Pereyra G, Schuldt B, Jansen S, Kattge J, Landhäusser SM, Levick SR, Preisler Y et al. 2017. A synthesis of tree functional traits related to drought-induced mortality in forests across climatic zones. *Journal of Applied Ecology.* doi: 10.1111/1365-2664.12874.
- Olmo M, Lopez-Iglesias B, Villar R. 2014. Drought changes the structure and elemental composition of very fine roots in seedlings of ten woody tree species. Implications for a drier climate. *Plant and Soil* 384: 113–129.
- O'Neill GA, Hamman A, Wang T. 2008. Accounting for population variation improves estimates of the impact of climate change on species growth and distribution. *Journal of Applied Ecology* 45: 1040–1049.
- Orwig DA, Abrams MD. 1997. Variation in radial growth responses to drought among species, site, and canopy strata. Trees 11: 474–484.
- Pan J, Wang B, Pei Z-Y, Zhao W, Gao J, Mao J-F, Wang X-R. 2015. Optimization of the genotyping-by-sequencing strategy for population genomic analysis in conifers. *Molecular Ecology Resources* 15: 711–722.
- Parent B, Hachez C, Redondo E, Simonneau T, Chaumont F, Tardieu F. 2009.

 Drought and abscisic acid effects on aquaporin content translate into changes in



- hydraulic conductivity and leaf growth rate: a trans-scale approach. *Plant Physiology* **149**: 2000–2012.
- Perdiguero P, Barbero MC, Cervera MT, Soto Á, Collada C. 2012a. Novel conserved segments are associated with differential expression patterns for Pinaceae dehydrins. *Planta* 236: 1863–1874.
- Perdiguero P, Collada C, Barbero Mdel C, García Casado G, Cervera MT, Soto A. 2012b. Identification of water stress genes in *Pinus pinaster* Ait. by controlled progressive stress and suppression-subtractive hybridization. *Plant Physiology and Biochemistry* 50: 44–53.
- Phillips RP, İbáñez I, D'Orangeville L, Hanson PJ, Ryan MG, McDowell NG. 2016. A belowground perspective on the drought sensitivity of forests: towards improved understanding and simulation. Forest Ecology and Management 380: 309–320
- Plaut JA, Wadsworth WD, Pangle R, Yepez EA, McDowell NG, Pockman WT. 2013. Reduced transpiration response to precipitation pulses precedes mortality in a Piñon–juniper woodland subject to prolonged drought. New Phytologist 200: 375–387.
- Plomion C, Bartholomé J, Bouffier L, Brendel O, Cochard H, de Miguel M, Delzon S, Gion J-M, Gonzalez-Martinez SC, Guehl J-M *et al.* 2016. Understanding the genetic bases of adaptation to soil water deficit in trees through the examination of water use efficiency and cavitation resistance: maritime pine as a case study. *Journal of Plant Hydraulics* 3: e008.
- Poland JA, Rife TW. 2012. Genotyping-by-sequencing for plant breeding and genetics. *Plant Genome* 5: 92–102.
- Prasolova NV, Xu Z, Farquhar GD, Saffigna PG, Dieters MJ. 2001. Canopy carbon and oxygen isotope composition of 9-year-old hoop pine families in relation to seedling carbon isotope composition, growth, field growth performance, and canopy nitrogen concentration. Canadian Journal of Forest Research 31: 673–681.
- Prunier J, Laroche J, Beaulieu J, Bousquet J. 2011. Scanning the genome for gene SNPs related to climate adaptation and estimating selection at the molecular level in boreal black spruce. *Molecular Ecology* 20: 1702–1716.
- Rehfeldt GE. 1991. A model of genetic variation for *Pinus ponderosa* in the inland northwest (USA): applications in gene resource management. *Canadian Journal of Forest Research* 21: 1491–1500.
- Rehfeldt GE, Hoff RJ, Steinhoff RJ. 1984. Geographic patterns of genetic variation in *Pinus monticola. Botanical Gazette* 145: 229–239.
- Rehfeldt GE, Leites LP, St Clair JB, Jaquish BC, Sáenz-Romero C, López-Upton J, Joyce DG. 2014. Comparative genetic responses to climate in the varieties of *Pinus ponderosa* and *Pseudotsuga menziesii*: clines in growth potential. *Forest Ecology and Management* 324: 138–146.
- Rehfeldt GE, Tchebakova NM, Parfenova YI, Wykoff WR, Kuzmina NA, Milyutin LI. 2002. Intraspecific responses to climate in *Pinus sylvestris. Global Change Biology* 8: 912–929.
- Reinhardt K, Germino MJ, Kueppers LM, Domec J-C, Mitton J. 2015. Linking carbon and water relations to drought-induced mortality in *Pinus flexilis* seedlings. *Tree Physiology* 35: 771–782.
- Resende MFR, Muñoz P, Acosta JJ, Peter GF, Davis JM, Grattapaglia D, Resende MDV, Kirst M. 2012. Accelerating the domestication of trees using genomic selection: accuracy of prediction models across ages and environments. *New Phytologist* 193: 617–624.
- Ripullone F, Lauteri M, Grassi G, Amato M, Borghetti M. 2004. Variation in nitrogen supply changes water-use efficiency of *Pseudotsuga menziesii* and *Populus* × *euroamericana*; a comparison of three approaches to determine water-use efficiency. *Tree Physiology* 24: 671–679.
- Ryan MG. 2011. Tree responses to drought. *Tree Physiology* 31: 237–239.
- Sala A, Piper F, Hoch G. 2010. Physiological mechanisms of drought-induced tree mortality are far from being resolved. *New Phytologist* 186: 274–281.
- Santos-Del-Blanco L, Bonser SP, Valladares F, Chambel MR, Climent J. 2013.
 Plasticity in reproduction and growth among 52 range-wide populations of a Mediterranean conifer: adaptive responses to environmental stress. *Journal of Evolutionary Biology* 26: 1912–1924.
- Schmidtling RC. 1994. Use of provenance tests to predict response to climatic change: loblolly pine and Norway spruce. *Tree Physiology* 14: 805–817.

- Seibt U, Rajabi A, Griffiths H, Berry JA. 2008. Carbon isotopes and water use efficiency: sense and sensitivity. *Oecologia* 155: 441–454.
- Sevanto S, McDowell NG, Dickman LT, Pangle R, Pockman WT. 2014. How do trees die? A test of the hydraulic failure and carbon starvation hypotheses. *Plant, Cell & Environment* 37: 153–161.
- Silim S, Guy R, Patterson T, Livingston N. 2001. Plasticity in water-use efficiency of *Picea sitchensis*, *P. glauca* and their natural hybrids. *Oecologia* 128: 317–325.
- Sperry JS, Donnelly JR, Tyree MT. 1988. A method for measuring hydraulic conductivity and embolism in xylem. *Plant, Cell & Environment* 11: 35–40.
- Sperry JS, Hacke UG, Pittermann J. 2006. Size and function in conifer tracheids and angiosperm vessels. American Journal of Botany 93: 1490–1500.
- Sperry JS, Nichols KL, Sullivan JEM, Eastlack SE. 1994. Xylem embolism in ringporous, diffuse-porous, and coniferous trees of northern Utah and interior Alaska. *Ecology* 75: 1736–1752.
- Sthultz CM, Gehring CA, Whitham TG. 2009. Deadly combination of genes and drought: increased mortality of herbivore-resistant trees in a foundation species. *Global Change Biology* 15: 1949–1961.
- Tardieu F, Simonneau T. 1998. Variability among species of stomatal control under fluctuating soil water status and evaporative demand: modelling isohydric and anisohydric behaviours. *Journal of Experimental Botany* 49: 419–432.
- Tyree MT, Ewers FW. 1991. The hydraulic architecture of trees and other woody plants. *New Phytologist* 119: 345–360.
- Valliyodan B, Nguyen HT. 2006. Understanding regulatory networks and engineering for enhanced drought tolerance in plants. Current Opinion in Plant Biology 9: 189–195.
- Velasco-Conde T, Yakovlev I, Majada JP, Aranda I, Johnsen Ø. 2012. Dehydrins in maritime pine (*Pinus pinaster*) and their expression related to drought stress response. *Tree Genetics & Genomes* 8: 957–973.
- Villar-Salvador P, Ocaña L, Peñuelas J, Carrasco I. 1999. Effect of water stress conditioning on the water relations, root growth capacity, and the nitrogen and non-structural carbohydrate concentration of *Pinus halepensis* Mill. (Aleppo pine) seedlings. *Annals of Forest Science* 56: 459–465.
- Virlouvet L, Fromm M. 2015. Physiological and transcriptional memory in guard cells during repetitive dehydration stress. New Phytologist 205: 596–607.
- Voltas J, Chambel MR, Prada MA, Ferrio JP. 2008. Climate-related variability in carbon and oxygen stable isotopes among populations of Aleppo pine grown in common-garden tests. *Trees* 22: 759–769.
- Wang T, Hamann A, Yanchuk A, O'Neill GA, Aitken SN. 2006. Use of response functions in selecting lodgepole pine populations for future climates. *Global Change Biology* 12: 2404–2416.
- Wang Z, Gerstein M, Snyder M. 2009. RNA-Seq: a revolutionary tool for transcriptomics. *Nature Reviews Genetics* 10: 57–63.
- Watkinson JI, Sioson AA, Vasquez-Robinet C, Shukla M, Kumar D, Ellis M, Heath LS, Ramakrishnan N, Chevone B, Watson LT et al. 2003. Photosynthetic acclimation is reflected in specific patterns of gene expression in drought-stressed loblolly pine. Plant Physiology 133: 1702–1716.
- Whitlock MC, Lotterhos KE. 2015. Reliable detection of loci responsible for local adaptation: inference of a null model through trimming the distribution of F_{ST} . The American Naturalist 186: S24–S36.
- Wyckoff PH, Clark JS. 2002. The relationship between growth and mortality for seven co-occurring tree species in the southern Appalachian mountains. *Journal of Ecology* 90: 604–615.
- Xu J, Lu J, Evans R, Downes GM. 2014. Relationship between ring width and tracheid characteristics in *Picea crassifolia*: implication in dendroclimatology. *BioResources* 9: 2203–2213.
- Yeaman S, Hodgins KA, Suren H, Nurkowski KA, Rieseberg LH, Holliday JA, Aitken SN. 2014. Conservation and divergence of gene expression plasticity following c. 140 million years of evolution in lodgepole pine (*Pinus contorta*) and interior spruce (*Picea glauca* × *Picea engelmannii*). New Phytologist 203: 578–591.
- Yuxiu Z, Zi W, Jin X. 2007. Molecular mechanism of dehydrin in response to environmental stress in plants. *Progress in Natural Science* 17: 237–246.
- Zhang JW, Feng Z, Cregg BM, Schumann CM. 1997. Carbon isotopic composition, gas exchange, and growth of three populations of ponderosa pine differing in drought tolerance. *Tree Physiology* 17: 461–466.