

REVIEW PAPER

The plant model system *Arabidopsis* set in an evolutionary, systematic, and spatio-temporal context

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Abstract

Arabidopsis thaliana is one of the prevailing plant model systems used for most facets of plant molecular, cell, and evolutionary biology. There are good reasons for that, such as easy cultivation, short generation time, a selfing reproductive system, and a wide geographic distribution with easy access to respective germplasm material. For the last two decades, the entire genus Arabidopsis with its various species has emerged as a model system allowing the study of additional traits and characters not found in A. thaliana. These species grow naturally under very different environmental conditions and mostly underwent independent evolutionary histories. Accordingly, consideration of their respective biogeographic and phylogenetic (taxonomic) context is important for any study aiming to explore fully the potential of comparative studies. Here, we review current understanding of spatio-temporal patterns of Arabidopsis evolutionary history and introduce the various taxa in more detail, including those traits well studied by the scientific community. The significance of polyploidization and interspecies gene flow is also discussed.

Keywords: Arabidopsis, biogeography, evolutionary history, phylogeny, speciation, taxonomy, trait evolution.

Introduction

The taxonomy of Arabidopsis as an emerging study and model system

The genus Arabidopsis, and in particular Arabidopsis thaliana (Brassicaceae), is among the few prevailing plant model species for studying fundamental biological processes from molecule to population level (Weigel, 2012; Provart et al., 2016). An excellent historical overview of how A. thaliana found its way into thousands of labs of plant molecular biologists has been provided by Koornneef and Meinke (2010). Its career as a model system started with Laibach's work using the species as a genetic system (Laibach, 1943), followed by renewed interest in a 'renaissance period' in the late 1970s and 1980s (reviewed in Meyerowitz, 2001; Koornneef and Meinke, 2010), and

accumulated into >55 000 indexed papers having *Arabidopsis* in the title, abstract, or keywords (Provart *et al.*, 2016).

During most of this time, systematics and the evolutionary history of the genus *Arabidopsis* were not an issue and they were not much taken into account until recently. Consequently, there is a parallel and often confusing taxonomic history regarding species and genus delimitations. Here, a benchmark was set upon entering the new millennium when a comprehensive new circumscription of the entire genus was introduced (reviewed in Koch *et al.*, 2008). This had some serious consequences for >60 *Arabidopsis* species (excluding *A. thaliana*) recognized at that time, most of which have been transferred to very different genera (Al-Shehbaz *et al.*, 1999). On

the other hand, the newly circumscribed genus Arabidopsis gained species because species shown by molecular markers to be the closest relatives of A. thaliana (O'Kane and Al-Shehbaz, 2003; Koch et al., 2008) were incorporated. This taxonomic history—changes of names on the genus and species level following distinct rules provided by the 'Code of nomenclature' (Turland et al., 2018)—is recapitulated here in in some detail, and it should be kept in mind that previous work published under species names such as Arabidopsis pumila, Arabidopsis griffithiana, or Arabidopsis wallichii (to name just a few) is in fact focused on species that are only distantly related to A. thaliana, and belong to different genera such as Olimarabidopsis or Crucihimalya (Kiefer et al., 2014; Koch et al., 2018). The confusion that can arise from assuming close relatedness based on genus names alone emphasizes the fact that reliable taxonomy and systematics are key to the interpretation of biological, especially evolutionary, information (Koch and German, 2013; Koch et al., 2018).

The taxonomic confusion surrounding A. thaliana started soon after 1872. Many species were transferred into a genus Arabidopsis based on a few simple morphological characters (latiseptate siliques terete to quadrangular in cross section; and branched trichomes) and, therefore, some 60 species were recognized in Arabidopsis (German and Ebel, 2005). With the pioneering work of O'Kane and Al-Shehbaz (1997) and Al-Shehbaz et al. (1999), this concept was completely changed and nine 'new' Arabidopsis species with several subspecies, mostly from the former genus Cardaminopsis, were recognized together with A. thaliana. This taxonomic revision was based on initial phylogenetic analysis using the nuclear-encoded internal transcribed spacer of rDNA (O'Kane et al., 1996) and later confirmed with a broader taxon sampling (O'Kane and Al-Shehbaz, 2003). It has to be noted that earlier, Hedge (1968) proposed a closer relationship between Arabidopsis and Arabis, while recognizing that the two genera differ only in the position of the cotyledons relative to the radicle in the seeds. Meanwhile we know from a series of phylogenetic studies that Arabis is also not monophyletic, and it was at that time that species from the whole Brassicaceae family (Karl and Koch, 2013) were resampled. It is curious to see that Hedge (1968) claimed that the Himalayan species A. wallichii (now Crucihimalya wallichii) is essentially an intermediate between the two genera. However, as molecular data were not available, Hedge and others could not have been aware that, in fact, the genus Arabis at that time was comprised of various unrelated evolutionary lineages (see Koch et al., 1999, 2000, 2001; Bailey et al., 2006). Nowadays the historical concept of a monophyletic genus Arabis no longer exists (Karl and Koch, 2013). Because taxa from both genera, Arabidopsis and Arabis, share this taxonomic history, they frequently exhibit related taxonomic nomenclatural problems with numerous misleading phylogenetic implications.

In a review entitled 'Arabidopsis and its poorly known relatives', it is >10 years ago that the closest relatives of A. thaliana were introduced in more detail (Clauss and Koch, 2006), with some particular focus on Arabidopsis lyrata, with >600 papers published using this species between 2000 and 2018, and >1350 papers published between 2000 and 2018 dealing with any of the 'true wild relatives' (Web of Science, accessed 7 July 2018).

The new concept of the genus Arabidopsis was presented 15 years ago (O'Kane and Al-Shehbaz, 1997, 2003) paralleled by a contribution from Koch et al. (1999) in which some taxonomical problems including both Arabis and Arabidopsis have been unravelled. Some species and subspecies were added later, either because of a transfer of taxa previously never associated with Arabidopsis (Warwick et al., 2006) or because of the description of a new species (Kadota, 2007). Changes since 1997 were mostly due to changing the rank of taxa as subspecies or recognizing synonyms (Dorofeyev, 2002; Marhold et al., 2003; Shimizu et al., 2005; Kolník and Marhold, 2006; Kadota, 2007; Elven and Murray, 2008). As a result, depending on the applied species concept, Arabidopsis can be considered as a genus comprising at least nine species and six subspecies (O'Kane and Al-Shehbaz, 1997); additionally, 15 species and 16 subspecies are presented with the latest and most up-to-date taxonomic status report (e.g. summarized in Koch et al., 2008; Hohmann et al., 2014) (Table 1).

An introduction to the biological system

The phylogenetic context

The genus Arabidopsis belongs to tribe Camelineae, and among its closest sister groups is the genus Capsella. The name is derived from Arabis (a genus of Brassicaceae, from which it cannot be distinguished by any unique character) and Greek opsis, in reference to this similarity. Members are mostly perennial and rarely annual or biennial. Multicellular glands are absent and trichomes are simple and mixed with stalked 1-3(or 4)-forked trichomes. Petals are white, pink, or purple, and nectar glands are confluent and subtending bases of all stamens. Fruits are dehiscent and formed as capsular, linear siliques. Other closely related genera are Neslia and Camelina. All three of them, Capsella, Neslia, and Camelina, are characterized by silicules. Recent phylogenetic data indicated incongruencies among gene trees comparing plastome and nuclear genome data and questioning the monophyly of tribe Camelineae and the sister relationship of Arabidopsis to a clade consisting of Neslia-Camelina-Capsella (Hohmann et al., 2015; Huang et al., 2016). However, Beilstein and co-workers have shown that the Arabidopsis versus Neslia-Camelina-Capsella relationship is the most likely sister relationship scenario, and this might be masked because of ancient gene flow among lineages predating most of the species considered (Forsythe et al., 2017, Preprint). In our current circumscription, tribe Camelineae is species poor (~50 species) (Koch et al., 2018), comprised of nearly 50% of neopolyploids, and its average 1 Cx (monoploid) genome size is ~0.36 pg (equals 300 Mbp, compared with 0.16 pg in A. thaliana representing ~135 Mbp) (Hohmann et al., 2015). The base chromosome number of Arabidopsis is n=8, with A. thaliana showing a derived situation with n=5chromosomes (Table 1). Within the genus Arabidopsis, phylogenetic analyses have recognized seven major lineages, with all of them also comprising diploid taxa: Arabidopsis halleri lineage, Arabidopsis lyrata lineage, Arabidopsis arenosa lineage, Arabidopsis pedemontana, Arabidopsis cebennensis, Arabidopsis croatica, and A. thaliana (Koch and Matschinger, 2007;

Table 1. The genus Arabidopsis and its various taxa

Arabidopsis thaliana L.		2x	Native range almost all Europe to Central Asia
Arabidopsis arenosa group			
A. arenosa (L.) Lawalrée ^a		$(2x) \ 4x$	
	subsp. borbasii (Zapał.) O'Kane &		Mountain ranges in Central and Western Europe
	Al-Shehbaz		
	subsp. arenosa		Central and Western Europe, Scandinavia at lower altitudes
A. carpatica nom. prov.		2 <i>x</i>	Carpathian Mountains in mountain ranges on calcareous bedrock
A. neglecta (Schult.) O'Kane & Al-Shehbaz			Carpathian Mountains
	subsp. robusta nom. prov.	4 <i>x</i>	Alpine regions and only occasionally at lower altitudes
	subsp. neglecta nom. prov.	2x	Alpine regions on siliceous bedrock types
A. nitida nom. prov.		2 <i>x</i>	Carpathian Mountains
A. petrogena (A. Kern.) V.I. Dorof.			Carpathian Mountains
	subsp. petrogena nom. prov.	2 <i>x</i>	Carpathians and Pannonian lowland
	subsp. exoleta nom. prov.	4 <i>x</i>	Carpathians at lower altitudes
Arabidopsis halleri group		2 <i>x</i>	
A. halleri (L.) O'Kane & Al-Shehbaz			
	subsp. dacica (Heuff.) Kolník		Carpathians, Romania
	subsp. gemmifera (Matsum.) O'Kane &		Russia Far East, NE China, Korea, Japan, and Taiwan
	Al-Shehbaz		
	subsp. <i>halleri</i>		Europe
	subsp. ovirensis (Wulfen) O'Kane &		Austria (taxon and its distribution is unclear)
	Al-Shehbaz		
	subsp. tatrica (Pawł.) Kolník		Tatra mountains, Slovakia
A. umezawana Kadota			Japan, Hokkaido (alpine zone of Mt. Rishirizin)
Arabidopsis lyrata group			
A. lyrata (L.) O'Kane & Al-Shehbaz			
	subsp. lyrata	2x (4x)	Alaska, Canada, USA
	subsp. petraea (L.) O'Kane &	2 <i>x</i>	Europe
	Al-Shehbaz		
	subsp. umbrosa nom. prov.	2 <i>x</i>	Arctic NE Asia, Siberia, Alaska, Canada
	subsp. septentrionalis nom. prov.	4x	Arctic NE Europe, European Russia to Siberia (may be admixed
			populations between subsp. lyrata and umbrosa)
A. arenicola (Richardson) Al-Shehbaz et al.		2 <i>x</i>	Arctic Canada and Greenland
Endemic diploid taxa			
A. croatica (Schott) O'Kane & Al-Shehbaz		2 <i>x</i>	Bosnia, Croatia
A. cebennensis (DC.) O'Kane & Al-Shehbaz		2 <i>x</i>	SE France
A. pedemontana (Boiss.) O'Kane &		2 <i>x</i>	NW Italy
Al-Shehbaz			
Allopolyploid taxa			
A. suecica (Fr.) Norrl.		4 <i>x</i>	Fennoscandinavia and the Baltic region
A. kamchatica (DC.) K. Shimizu & Kudoh		4 <i>x</i>	Boreal Alaska, Canada, E Siberia, Russian Far East, Korea, Japan
subsp. kamchatica			Taiwan
A. kamchatica subsp. kawasakiana (Makino)		4 <i>x</i>	Japan, winter annual (coastal, lowland)
K. Shimizu & Kudoh			

Species-level taxonomy refers to Al-Shehbaz et al. (1999), Koch et al. (2008), and Hohmann et al. (2014). The most recent treatment of subspecies within A. halleri has been proposed by Kolnik and Marhold (2006). Arabidopsis arenicola has been introduced by Warwick et al. (2006) on the species level. Ploidal levels (diploid=2x, tetraploid=4x) are summarized from Hohmann et al. (2014) and Kolár et al. (2016b). Base chromosome number is n=8 (2n=2x=16), with A. suecica (2n=4x=26) and A. thaliana (2n=2x=10) as the only exceptions.

Hohmann et al., 2014; Novikova et al., 2016) (Fig. 1, Table 1). Although substantial ancient [million(s) of years ago] and contemporary gene flow (past and interglacial) has been demonstrated (e.g. Jørgensen et al., 2011; Arnold et al., 2015; Novikova et al., 2016; Hohmann and Koch, 2017), these major lineages are well defined, genetically, ecologically, and taxonomically. Also it becomes evident that this source of natural variation including A. thaliana itself may serve as an excellent system to

study evolutionary processes and ecological strategies within an environmental context (reviewed for A. thaliana by Takou et al., 2018).

Species and subspecies in Arabidopsis are taxonomic units and are defined by morphological descriptions including information on cytology, ecology, or distribution. Accordingly, any validly described taxon—which refers to species but also to subspecies and varieties—following the rules of botanical

^a There is another variety still treated under the old genus name Cardaminopsis: C. arenosa var. intermedia (Neilr.) Hayek from high alpine regions in the south-eastern Austrian Alps (Seckauer Tauern).

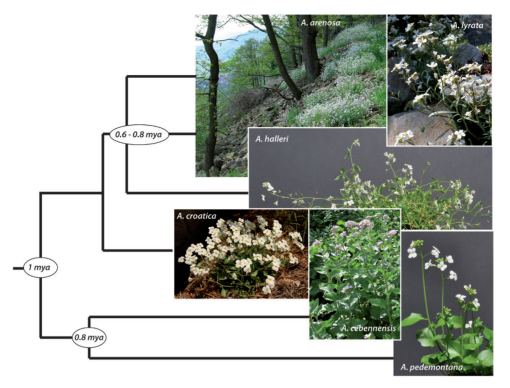


Fig. 1. Schematic phylogenetic relationships among lineages of relatives of Arabidopsis thaliana. Arabidopsis thaliana split from this entire group during Early Pliocene/Late Miocene ~5.8 million years ago (mya). Divergence times are summarized from Hohmann et al. (2015), Arnold et al. (2015), Hohmann (2016), and Hohmann and Koch (2017).

nomenclature can serve as accepted taxon. As a consequence, there is no general agreement or taxonomic concept when a subspecies should be considered on species rank or vice versa. Obviously, this needs to be kept in mind for the genus Arabidopsis, too. In some lineages, such as A. halleri, taxonomic subgrouping has mostly been introduced on a subspecies level, but in A. arenosa taxomomic grouping of the entire complex was done with preference for a species-level concept. However, most important is that the taxa are defined, on either a species or a subspecies level, with unambiguous names linked with all information on the respective taxon (Koch et al., 2018)

Arabidopsis halleri: metals and soils

Arabidopsis halleri is a perennial, outcrossing, diploid, creeping species and its various subspecies are widely distributed in Europe and Eastern Asia. Subspecies definition is often based on morphology, ecology, and distribution, and a discussion of initial evidence for correlation with genetic data is summarized below. A first draft genome was recently published (Briskine et al., 2017). This species grows on metal-rich soils and exhibits high accumulation of zinc, cadmium, and lead. The species is one among few which are heavy metal hypertolerant (Ernst, 1974) and are able to colonize metalliferous soils. Furthermore, the species also shows the rare trait of metal hyperaccumulation (e.g. Hanikenne et al., 2008, and references therein), and it has been shown that the complex interactions of plants with contaminated soils are not only environment dependent but also geographically structured (Stein et al., 2017). However, the entire distribution range in montane to alpine environments is also characterized by non-metalliferous soils, and large-scale phylogeographic analysis and ecological niche modelling indicate complex post-glacial recolonization scenarios (Wasowicz et al., 2016). Since metallicolous populations have been founded separately all over the distribution range, this trait seems to have arisen multiple times from distinct populations that do not exhibit that trait (Pauwels et al., 2005). Among the other Arabidopsis species, initial work on bedrock adaptation has been done on North American A. lyrata from serpentine soils (Turner et al., 2008, 2010), or A. arenosa from serpentine stands in Eastern Austria (Arnold et al., 2016).

Within A. halleri, up to five subspecies have been recognized (for details, refer to Koch et al., 2008) with two of them, A. halleri subsp. gemmifera and A. halleri subsp. ovirensis, accepted by some authors as separate species, namely A. gemmifera and A. ovirensis, respectively (Iljinska et al., 2007; Kadota, 2007). In Central Europe three subspecies can be differentiated (Kolnik and Marhold, 2006). The most widely distributed taxon is A. halleri subsp. halleri found from Poland to Serbia and from France to Western Ukraine. The second taxon, A.halleri subsp. tatrica, is a Western Carpathian endemic species. In contrast to these two subspecies, which are substrate indifferent, the third subspecies, A. halleri subsp. dacica, from the Eastern and Southern Carpathians is an alpine and rarely montane species from acidic bedrock types. The Eastern Asian subspecies A. halleri subsp. gemmifera is geographically and genetically fully isolated (Koch et al., 2008); and A. halleri subsp. ovirensis described as endemic to the East Austrian high mountain range at Mount Obir, in Carinthia, might not represent a genetically defined taxon but rather an isolated population from within the A. halleri subsp. halleri gene pool (Koch and Matschinger, 2007; Koch et al., 2008). There are some first phylogeographic studies, and

it has been shown that one major centre of genetic diversity is located in the Eastern Austrian Alps (Koch and Matschinger, 2007; Pauwels et al., 2012).

A comprehensive phylogeographic scenario (Pauwels et al., 2012) also demonstrated high levels of genetic diversity found in the eastern region of the European Alps and was explained convincingly by admixture and secondary contact of different European gene pools. The evolutionary scenario of A. halleri is best placed among Pleistocene glaciation and deglaciation cycles (Koch and Matschinger, 2007; Wasowicz et al, 2016). In a detailed evolutionary study, Roux et al. (2011) suggested the onset of radiation within A. halleri subsp. halleri to be ~335 000 years ago but, because this study does not consider other subspecies, a much deeper evolutionary split is likely, as indicated more recently (Novikova et al., 2016). Generally, the number of genetic-evolutionary studies centred around A. halleri is high (e.g. Van Rossum et al., 2004; Meyer et al., 2009; Heidel et al., 2010), and various ecological studies consider traits such as herbivory (Kawagoe and Kudoh, 2010; Sato and Kudoh, 2017), flowering time (Shimizu et al., 2011), or reproduction (Llaurens et al., 2008).

Arabidopsis lyrata: breeding system and evolutionary biology

Arabidopsis lyrata is a perennial and mostly outbreeding (selfincombatible) species with five taxa in total. Its close relationship to A. thaliana and a genetically diverse sporophytic self-incompatibility system fostered its early establishment as a model system to study breeding system evolution and its genetic and molecular components (e.g. Nasrallah et al., 2004; Guo et al., 2011; Vekemans et al., 2014). Various aspects of trait evolution in a perennial plant have also been studied intensively, such as thermal stress response (Wos and Willi, 2018), seed biology (Hämälä et al., 2017), or general fitness and survival (Leinonen et al., 2011). More details and earlier ecological work have been summarized by Clauss and Koch (2006). The species is one of the model systems used to study local adaptation in the context of trait evolution. One of those traits is adaptation in response to heavy metal such as nickel, thereby linking this species with A. halleri (e.g. Turner et al., 2008, 2010; Veatch-Blohm et al., 2017); other traits are herbivory response (Vergeer and Kunin, 2011), flowering time (Hämälä et al., 2018), or more generally pathogen-mediated selection (Buckley et al., 2018). Studies of molecular evolution have been fostered by the publication of the A. lyrata genome (Hu et al., 2011) and have focused on various features including transposons (Bonchev and Willi, 2018), transcriptional changes (Wos and Willi, 2018), or accumulation of mutational load (Willi et al., 2018). Research on populations in North America at the Great Lakes shows that some of them are undergoing a loss of a functional self-incompatible (SI) system (Mable et al., 2005), which allows the study of causes and genomic consequences of mating shifts (e.g. Mable et al., 2005, 2017; Griffin and Willi, 2014) and to compare those with largely inbreeding A. thaliana. Artifical interspecific hybrids of A. lyrata with Arabidopsis were created to study the expression pattern of transposable elements (TEs) in the hybrid genomic context and showing that TE expression is globally robust (Göbel et al., 2018). An artificial interspecific hybrid of A. lyrata with A. halleri elucidated the role of cis-acting modifications in cold adaption (He et al., 2016)

The rosette-forming plant has a circumpolar distribution. The two main distribution centres are formed by subsp. petraea in Central and Northern Europe and subsp. lyrata in Northern America. This disjunct distribution is mediated by subsp. umbrosa and subsp. septentrionalis (Table 1). The fifth taxon, Arabidopsis arenicola, was originally described as Arabis and later transferred to Arabidopsis as a separate species. Arabidopsis arenicola is an arctic and self-compatible (MAK, unpublished results) species from Canada and Greenland of very recent and probably post-glacial origin from subsp. lyrata (Hohmann et al., 2014). Therefore, it might be best treated in future as another subspecies of A. lyrata. The species complex is diploid with two exceptions: A. lyrata subsp. sepentrionalis probably represents a tetraploid taxon; it might have arisen via secondary contact between the two main subsp. lyrata and subsp. petraea gene pools and may represent an 'allo'tetraploid (Hohmann et al., 2014). The other exceptions are tetraploid populations from the northeastern limestone Forealps and pannonical dryland regions close to the Danube river in Austria. Those originated from adaptive introgression between two species (A. lyrata and A. arenosa) and are discussed separately below (Schmickl and Koch, 2011; Hohmann and Koch, 2017).

Studying six populations, Ross-Ibara et al. (2008) presented an estimate of the split time between the European and North American distribution ranges of ~35 000 years based on demographic models and assuming a mutation rate of 1.5×10^{-8} (following Koch et al., 2000; see also the section on A. arenosa for application of different mutational rates). This was also compared with $F_{\rm ST}$ -based comparisons which result instead in an estimate of ~90 000-170 000 years ago. A recent study estimated a split time between subsp. petraea and subsp. lyrata at 190 000-310 000 years ago (Mattila et al., 2017). This much better fits presented comprehensive phylogeographic scenarios covering the entire distribution range. It also considers the amphiberingean region (which means that the range is centred on what was formerly Beringia and that the species occurs on both sides of the Bering Strait) as a melting pot of lineages coming into secondary contact during the last glaciation and deglaciation cycles (e.g. Schmickl et al., 2008, 2010; Hohmann et al., 2014).

A phylogeographic study focusing on North American subsp. lyrata was not able to show any clear structure of genetic variation (Tedder et al., 2010), tentatively because of a very recent post-glacial biogeography. Similarly, Hämälä et al. (2018) demonstrated post-glacial evolutionary demographic scenarios in Scandinavia. Central European populations might have diverged from direct ancestors of Scandinavian populations ~28 000 generations ago. This equals ~56 000 years ago considering the life cycle of this plant which flowers latest in its second year.

Arabidopsis arenosa: polyploid evolution and interspecies gene flow

The A. arenosa complex is definitely the most complex and diverse group of taxa. Although its overall distribution is smaller than that of A. lyrata and A. halleri, the taxa occupy a wide range of habitats from alpine regions to sandy coastal sites and dune vegetation. The species complex consists of short-lived perennials, and the original rosette often dies after flowering. Daughter rosettes are frequently formed. At the moment, five species with various subspecies (Table 1) are recognized. However, in various cases, the respective names are provisional, such as A. nitida. There are tetraploid and diploid taxa, and it was only recently that a single origin of the tetraploids was demonstrated (Arnold et al., 2015). This may be surprising since a large-scale genetic screen using microsatellite variation demonstrated that tetraploids are genetically very diverse and mask the primary centre of diversity of diploids in the Carpathians (Schmickl et al., 2012). One explanation could be that tetraploid A. arenosa and its tetraploid subspecies originated from within large populations carrying a substantial amount of genetic variation. Furthermore, at least some interand intraspecies level gene flow might have occurred after polyploidization, even crossing ploidy levels (Jørgensen et al., 2011; Arnold et al., 2015) and thereby contributing substantial genetic variation. Intensive genetic interplay of A. arenosa has also been demonstrated with A. lyrata (Schmickl and Koch, 2011) and A. halleri (Hohmann, 2011; Hohmann et al., 2014) (Fig. 2).

Arabidopsis arenosa contributed as paternal hybridization partner to give rise to A. suecica (Koch and Matschinger, 2007; Novikova et al., 2017). Meanwhile A. suecica served as a model for interspecies A. thaliana hybrids (Säll et al., 2003), and studies using artificial hybrids between A. arenosa and Arabidopsis have been used (Nasrallah et al., 2000) to study gene silencing (e.g. Comai et al., 2000) or post-zygotic hybridization barriers (Bushell et al., 2003).

More recently, with Yant et al. (2013) and the overview given by Bomblies and Madlung (2014), A. arenosa was introduced as a system to study polyploidization. It also serves as a model system to study the molecular basis and evolution of meiosis in polyploids (e.g. Bomblies et al., 2015; Yant and Bomblies, 2017). This work also hints at interactive effects with important environmental factors such as temperature (Wright et al., 2015; Morgan et al., 2017).

Arabidopsis lyrata as well as A. arenosa only rarely occur on metalliferous soils, the exceptions being serpentine (e.g. in Austria), and it has been shown that parallel adaptation of different populations occurred via post-glacial adaptive introgression of genes from populations from the same species but also from A. lyrata (Arnold et al., 2016). This is another example of interspecies and interploidal gene flow since in this area A. arenosa is exclusively tetraploid and A. lyrata is diploid. The species split time between A. lyrata and A. arenosa was calculated as ~400 000 generations, which corresponds to ~800 000 years ago. This is in line with previous assumptions of a Mid- to Late-Pleistocene species split (Hohmann et al., 2014;

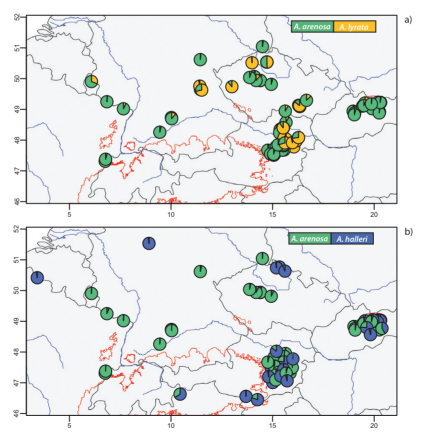


Fig. 2. Examples for genetic admixture between the three more widely distributed species groups in Central Europe. Genetic admixture is shown (a) for A. Iyrata-A. arenosa and (b) for A. halleri-A. arenosa. Genetic admixture data are taken and redrawn as pie charts from earlier published microsatellite data (Hohmann, 2011; Hohmann et al., 2014). (Genetic assignment analysis, number of genetic clusters: K=2 each, using Structure; Falush et al., 2007.) Glacial border from the Last Glacial Maximum (~25 000 years ago) are drawn as red lines based on Gibbard (2002) and Zasadni and Klapyta (2014).

Novikova et al., 2016). However, plastome data indicate a species divergence, which is younger and of ~550 000 years ago (Hohmann and Koch, 2017).

A single origin of all tetraploids in A. arenosa (Arnold et al., 2015) was estimated to ~11 000-33 000 generations ago and, considering phylogenetic relationships among tetraploid lineages, the centre of origin is best placed within the Carpathian Mountains. Geographically this matches the centre of species diversity of the A. arenosa lineage (Koch and Matschinger, 2007) as well as the distribution centre of diploids within the A. arenosa lineage (Kolár et al., 2016b). Extensive interglacial, glacial, or post-glacial range expansion of diploids has not been demonstrated (Kolár et al., 2016a), but a few diploids have migrated post-glacially towards the North into the Baltic. This also matches spatio-temporally with post-glacial hybrid speciation of Fennoscandinavian A. suecica (Novikova et al., 2017). However, it should be mentioned that the age estimate of tetraploid A. arenosa (Arnold et al., 2015) was based on a mutation rate of 3.7×10^{-8} . This is five times higher than the rates measured in A. thaliana in laboratory-based experiments (Ossowski et al., 2010); and applying those rates for the origin of tetraploid A. arenosa would result in a five times higher age estimate of ~110 000-330 000 years ago (compared with 11 000-33 000 generations with the 2 year life cycle equalling 22 000-66 000 years). This does correlate well with a spatiotemporal scenario of introgression of tetraploid A. arenosa into A. lyrata in Eastern Austria after the Riss glaciation (130 000 years ago) (Hohmann and Koch, 2017).

The diploid and local endemics: A. cebennensis, A. pedemontana, and A. croatica—from the French volcanic Massif Central to the Croatian Karst mountains

There are three species with a particularly narrow distribution. All of them are also phylogenetically separated from the other major lineages (Fig. 1). Arabidopsis cebennensis and A. pedemontana represent sister species and have the smallest distribution range in the whole genus (Koch et al., 2008). Arabidopsis cebennensis is restricted to the mountainous regions in Southern France at elevations ranging from 900 m to 1500 m above sea level. The highly disjunct populations occur in the Massif Central, the Cevennes, and the Ardeche regions. The outcrossing and perennial species is the tallest among all Arabidopsis relatives and can reach a size of up to 1 m. It grows in a narrow ecological niche in riverine systems often close to natural springs and characterized by continuously running cold mineral-rich water.

Arabidopsis pedemontana is restricted to a range of 50 km² in the Piedmont region of the north-western Italian Alps, at altitudes ranging from 1300 m to 2200 m above sea level. Also A. pedemontana is presumably self-incombatible, with a strong tendency for vegetative reproduction by clonal growth (Hohmann et al., 2014), but it resembles a growth type more similar to A. halleri (Fig. 1). Both species demonstrate very strong inter-regional genetic differentiation indicating severe genetic bottlenecks due to past glaciations (Jacquemin et al., 2016, Preprint). Drastic reductions of effective population

sizes have been demonstrated and placed those events with a maximum of 4000 and 40 000 years ago for A. cebennensis and A. pedemontana, respectively (Jacquemin et al., 2016, Preprint). Because the same authors questioned the dimension of the timing results considering their calculated and highly contradictory species split time of 160 000 years ago (see Fig. 1, compared ith 800 000 years ago), it is worth considering five times higher estimates. This would place the drastic bottleneck events for both species close to the glaciation maxima 25 000 years ago (Würm glaciation, A. cebennensis) and 130 000 years ago (Riss glaciation, A. pedemontana). Both species are genetically isolated from any other Arabidopsis species, and they also do not exhibit significant signatures of past gene flow among different species as was demonstrated for the other taxa (Koch and Matchinger, 2007; Hohmann et al., 2014; Novikova et al., 2016). However, ancient gene flow between the common ancestor of A. pedemontana and A. cebennensis with A. halleri from its western distribution range is nonetheless likely (Novikova et al., 2016).

Arabidopsis croatica is distributed along the Dinaric Mountains on rocky outcroppings and fissures in Croatia. The thermophylous diploid species is perennial and self-incompatible, and it is restricted to calcareous bedrock. Phylogenetic analyses placed this taxon as a sister species basal to A. lyrata and A. arenosa (Novikova et al., 2016; sequence data from re-sequenced nuclear genomes), and maternal signatures from entire plastome sequences even placed this taxon basal to all three species complexes (A. halleri, A. lyrata, and A. arenosa). However, only very few individuals of A. croatica have been analysed so far, and a more comprehensive sampling is needed to explain the incongruency which might reflect incomplete lineage sorting, secondary gene flow, or ancestral shared polymorphisms. It is interesting to see that genetic admixture analysis assuming five genetic clusters among perennial Arabidopsis relatives shows A. croatica sharing substantial genetic variation with A. cebennensis and A. pedemontana (Novikova et al., 2016), which cannot be explained by (unlikely) gene flow during the last 130 000 years and which may indeed indicate massive shared ancestral genetic variation.

All three species are endangered, and A. croatica and A. pedemontana are strictly protected according to national law and listed as 'critically endangered' according to the IUCN definition. Arabidopsis cebennensis is most often found in protected nature reserve areas, and thereby also receives full protection. Hence, major efforts should be undertaken to cultivate these species and provide germplasm resources, with the scientific community following the regulations of the Convention on Biological Diversity and the Nagoya protocol.

Allopolyploid systems: whole-genome duplications in A. kamchatica and A. suecica

There are two contrasting allopolyploid species systems, which have been nicely introduced in a recent review (Novikova et al., 2018), thereby highlighting the origin of these polyploid taxa around recent glaciation maxima. The Brassicaceae family as a whole is characterized by a high percentage of polyploid taxa. More than 43% of the species are neopolyploids (Hohmann

et al., 2015), and actually 10 mesopolyploidization events predating the evolution of tribes (monophyletic assemblages of genera) have been described (Lysak et al., 2005, 2007; Haudry et al., 2013; Mandákova et al., 2017). A family-wide summary of genome size and ploidy level evolution is provided in Hohmann et al. (2015) and indicates tremendous evolutionary dynamics of the genomes in size and structure, and continuous and rapid genome downsizing after polyploidization events (Lysak et al., 2009; Mandákova and Lysak, 2018). Therefore, polyploidization in Brassicaceae can be regarded as one of the driving mechanisms of adaptation to rapidly changing environments and is often able to explain the high speciation rates found (Jordon-Thaden et al., 2013). On the family level, there is also a significant correlation of lower 1 Cx (monoploid) genome size with annual life cycle in Arabidopsis (Hohmann et al., 2015) compared with its perennial wild relatives.

Arabidopsis suecica (2n=4x=26) had an origin with A. thaliana (2n=2x=10) as maternal partner providing an unreduced gamete and tetraploid A. arenosa providing the parental n=16 gamete. This plastid (i.e. maternally inherited) molecular signature was demonstrated more than two decades ago (Mummenhoff and Hurka, 1995; O'Kane et al., 1996). Chloroplast and nuclear markers suggest that A. suecica arose just once (Säll et al., 2003; Jakobsson et al., 2006). An analysis of haplotype variation in the A. thaliana portion of the A. suecica genome was recently used to argue that there were multiple origins (Novikova et al., 2016). The time of origin was estimated to be ~16 000 years ago in Eastern Europe or Central Eurasia, with subsequent migration into the Fennoscandinavian region while glaciers were retreating (Novikova et al., 2017). Whether it has a single or multiple origins, A. suecica is certainly of a more regional origin and carries limited genetic variation.

Showing high levels of genetic diversity compared with A. suecica, another allopolyploid, A. kamchatica (2n=4x=32), originated via hybridization and polyploidization of outcrossing diploid A. lyrata and diploid A. halleri subsp. gemmifera in Eastern Asia, presumably around Japan, north-eastern China, and eastern Siberia (Shimizu-Inatsugi et al., 2009; Schmickl et al., 2010). A post-glacial origin ~20 000 years ago is likely (Tsuchimatsu et al., 2012), but there is still some uncertainty because of different mutation rates applicable to the analyses. Both A. suecica and A. kamchatica are selfing species. In A. suecica, the self-incombatible system seems to have been overcome just once at the beginning of its evolutionary history (Novikova et al., 2017), whereas A. kamchatica underwent transitions to selfing several times via degradation of male specificity genes (Tsuchimatsu et al., 2012; Shimizu and Tsuchimatsu, 2015). Arabidopsis suecica is an annual plant like its maternal parent, A. thaliana, while the amphiberingean A. kamchatica subsp. kamchatica, which occurs on both sides of the Bering Strait, is a short-lived perennial, though in Japan the subspecies kawasakiana is also annual, in association with its adaptation to highly disturbed sandy seashore habitats at Lake Biwa in Japan (Higashi et al., 2012).

The combination of two different genomes in natural allopolyploids attracts research to study various aspects of genomic interactions. Both species allow study of the interplay of genomes which diverged nearly 6 million years ago

(A. suecica: A. thaliana versus A. arenosa) and ~600 000 years ago (A. kamchatica: A. lyrata versus A. halleri) with a similar time span of allopolyploid evolutionary history of ~20 000 years. These studies focus on gene expression (e.g. Tian et al., 2014; Miller et al., 2015; Paape et al., 2016), epigenetic regulation (Costa-Nunes et al., 2010; Pontvianne et al., 2012), retention of homologues (Chang et al., 2010; Paape et al., 2018, Preprint), or heterosis effects (Solhaug et al., 2016). Furthermore, ecological aspects of adaptation are addressed, such as cold tolerance (Armstrong et al., 2015), flowering biology (Kenta et al., 2011), trichome production and reproductive trade-off evolution (Steets et al., 2010), and also the ability to cope with heavy metal-rich soils (Paape et al., 2016) by retaining the ability to hyperaccumulate zinc from A. halleri.

Arabidopsis thaliana: evolutionary history beyond the 1001 genomes

Since the first phylogeographic studies on A. thaliana were published more than a decade ago (Sharbel et al., 2000; Beck et al., 2008), the 1001 Arabidopsis genomes consortium provided the first comprehensive insights into A. thaliana evolutionary history. These indicate glacial refuge areas on the Iberian Peninsula, the Canary and Cape Verde Islands, Sicily, and the Levant region (1001 Genomes Consortium, 2016). A very dynamic post-glacial spatio-temporal diversification pattern all over Eurasia was shown, with strong footprints of selection correlated with post-glacially changing climate and environment. This study highlighted the importance of the Last Glacial Maximum in structuring the present-day A. thaliana populations, and the detected spatio-temporal signature collapsed ~130 000 years ago, which exactly matches the preceding glacial maximum of the Mindel glacial. This highlights again, as shown for wild allopolyploid relatives (Novikova et al., 2018), that diversification processes in space and time are often set into the context of glacial maxima and the subsequent early phase of the generally shorter interglacial intervals. More detailed studies on A. thaliana (Brennan et al., 2014; Durvasula et al., 2017; Zou et al., 2017; Fulgione et al., 2018) elucidated multiple colonizations of the Iberian Peninsula via Central Europe and via Africa during the last glaciation (45 000 and 90 000 years ago, respectively) but pre-dating the Last Glacial Maximum, and large-scale range expansions in Africa at ~120 000-90 000 years ago were demonstrated. Africa was shown to be one primary cradle of genetic diversity (Durvasula et al., 2017) of present-day A. thaliana, with the transition to selfing occurring in North-western Africa. It was concluded that African populations best represent the early history of the species (Fulgione and Hancock, 2018). However, this conclusion does have some restrictions, because the presented modelling of nuclear genomic data does not reveal deeper evolutionary splits than 130 000 years ago and thereby points to the last common ancestor of present-day A. thaliana populations. Furthermore, there are no Eurasian A. thaliana populations older than any African lineages. However, there are at least a few indirect hints allowing tracing of even deeper evolutionary trajectories in A. thaliana. S-locus haplotype variation suggests a geographic localization of the transition to selfing in Africa

(Durvasula et al., 2017), and comparative studies among various Arabidopsis species estimated this transition to have occurred 413 000 years ago (Bechsgaard et al., 2006) or even ~1 million years ago (Tang et al., 2007). Assuming that these older estimates are reliable, the African history of A. thaliana is indeed nearly 1 million years old. However, A. thaliana split from its relatives >5 million years ago (e.g. Hohmann et al., 2015; Huang et al., 2016; Guo et al., 2017) and, therefore, more >4 million years of evolutionary history remains unclear.

It is worth mentioning that for several genera of Brassicaceae there is a phylogenetic sister relationship of a species-poor clade of annuals (often with one single species only) and a predominantly perennial clade. Environmental changes (e.g. increasing temperature or drought) led to the realization of two different strategies: (i) transition to an annual life cycle with later transition to selfing, rapid colonization of low-elevation habitats; and (ii) migration to the North and in particular towards higher elevations, thereby favouring niche differentiation, allopatric speciation, and increased speciation (Karl and Koch, 2013; Koch et al., 2016). Accordingly, it may be hypothesized that ~1 million years ago two perennial lineages evolved rapidly and independently. One lineage may have successfully colonized the African continent with a transition towards selfing and an annual life cycle. The other lineage underwent a complex evolutionary history leading to speciation in Eurasia and the amphiberingean region and leaving deep footprints of its development within its plastid genomes (Novikova et al., 2016; Hohmann and Koch, 2017). This scenario is in agreement with the finding of ancient gene flow between A. thaliana and A. lyrata but not between A. thaliana and A. arenosa or A. halleri (Novikova et al., 2016). However, most probably the detailed biogeographical context will remain a mystery, because we have no data for paleoenvironments from 1 million years ago and present distribution ranges do not allow comparative phylogeographic conclusions in deeper times.

The temporal context might be solved in much more detail in the future. Estimation of a temporally resolved evolutionary scenario largely depends on assumptions on the underlying mutational rate of DNA. Here it is important to distinguish between the mutation rate, which is the rate at which genomes change due to DNA damage, faulty repair, gene conversion, and replication errors, and the substitution rate, which is the rate at which mutations accumulate within a given gene pool (Exposito-Alonso et al., 2018). Therefore, it is important that any rate used is critically chosen, set into the right context, and compared carefully with other studies. The following examples apply for the nuclear genome. Studies spanning hundreds of millions of years and analysing various plant families including Brassicaceae used, for example, synonymous substitution rates of $6-7 \times 10^{-9}$ substitutions per site per year (Lynch, 2010; De la Torre et al., 2017), data sets focusing on the Brassicaceae family and spanning a few million years assumed $3.83-8.56 \times 10^{-9}$ (Huang et al., 2012) or 8.22×10^{-9} (Kagale et al., 2014) synonymous mutations per site per year, whereas in a study focusing on Capsella and Arabidopsis, a substitution rate per site per year of 7×10^{-9} was used (Wu et al., 2017). Other studies aiming to resolve time spans of a few hundreds or thousands years have elaborated substitution rates of 2.7×10^{-9} substitutions per site and generation for Arabidopsis (Exposito-Alonso et al., 2018), which was translated into $2-5 \times 10^{-9}$ substitutions per site per year as A. thaliana is an annual. A mutation rate of A. thaliana under 'lab conditions' was estimated and introduced with 7×10^{-9} mutations per site per year per generation (Ossowski et al., 2010). Finally, Arnold et al. (2015) simulated a rate for their analyses in A. arenosa of 3.7×10^{-8} substitutions per site per year. In summary, there is a great demand carefully to document and explain the nature of the chosen rates and compare results with those obtained using alternative rates and thereby allow cross-study comparisons.

Summary: bridging Arabidopsis thaliana and its relatives

This review highlighted that A. thaliana and its relatives do not just share a common ancestor at ~6 million years. After this split, the various species shared a spatio-temporal context to varying degrees. Severe environmental changes, such as via oscillating climate change during the entire Pleistocene, affected some of the lineages. This did not only lead to differing patterns and processes, but some are similar among them. This makes the genus Arabidopsis as a whole a suitable system to conduct comparative and across-species studies given that reliable spatio-temporal evolutionary scenarios are available. It is also summarized that secondary genetic contact is often triggered by glaciation maxima and subsequent shorter warming phases, and has affected any lineage including A. thaliana. There are many open questions such as the 'deep history' of A. thaliana or more detailed analyses differentiating between contemporary and past gene flow among A. lyrata, A. arenosa, and A. halleri. So far, most often comparative studies have involved only two species but, with the knowledge of putative gene flow among all species, future studies might consider multispecies approaches.

Comment on germplasm accessibility

It has to be highlighted that for some of the perennial relatives of Arabidopsis, access to natural populations requires not only considering local/regional regulations (e.g. protected areas such as nature reserves or national parks) and international regulations of benefit sharing (Convention of Biological Diversity, Nagoya protocol from 2014), but some species are actually under severe threat and are categorized according to IUCN protection definition categories as 'critically endangered', such as A. croatica and A. pedemontana. Others are locally endangered, and some populations of A. lyrata, in particular in Eastern Austria, are regionally endangered and may even be 'oversampled'.

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References

Al-Shehbaz IA, O'Kane SL, Price RA. 1999. Generic placement of species excluded from Arabidopsis. Novon 9, 296-307.

Armstrong JJ, Takebayashi N, Sformo T, Wolf DE. 2015. Cold tolerance in Arabidopsis kamchatica. American Journal of Botany 3, 439-448.

Arnold B, Kim ST, Bomblies K. 2015. Single geographic origin of a widespread autotetraploid Arabidopsis arenosa lineage followed by interploidy admixture. Molecular Biology and Evolution 32, 1382-1395.

Arnold BJ, Lahner B, DaCosta JM, Weisman CM, Hollister JD, Salt DE, Bomblies K, Yant L. 2016. Borrowed alleles and convergence in serpentine adaptation. Proceedings of the National Academy of Sciences, USA 113, 8320-8325.

Bailey CD, Koch MA, Mayer M, Mummenhoff K, O'Kane SL Jr, Warwick SI, Windham MD, Al-Shehbaz IA. 2006. Toward a global phylogeny of the Brassicaceae. Molecular Biology and Evolution 23, 2142-2160.

Beck JB, Schmuths H, Schaal BA. 2008. Native range genetic variation in Arabidopsis thaliana is strongly geographically structured and reflects Pleistocene glacial dynamics. Molecular Ecology 17, 902-915.

Bechsgaard JS, Castric V, Charlesworth D, Vekemans X, Schierup MH. 2006. The transition to self-compatibility in Arabidopsis thaliana and evolution within S-haplotypes over 10 Myr. Molecular Biology and Evolution 23, 1741-1750.

Bomblies K, Higgins JD, Yant L. 2015. Meiosis evolves: adaptation to external and internal environments. New Phytologist 208, 306-323.

Bomblies K, Madlung A. 2014. Polyploidy in the Arabidopsis genus. Chromosome Research 22, 117-134.

Bonchev G, Willi Y. 2018. Accumulation of transposable elements in selfing populations of Arabidopsis lyrata supports the ectopic recombination model of transposon evolution. New Phytologist 219, 767-778.

Brennan AC, Méndez-Vigo B, Haddioui A, Martínez-Zapater JM, Picó FX, Alonso-Blanco C. 2014. The genetic structure of Arabidopsis thaliana in the south-western Mediterranean range reveals a shared history between North Africa and southern Europe. BMC Plant Biology 14. 17.

Briskine RV, Paape T, Shimizu-Inatsugi R, Nishiyama T, Akama S, Sese J, Shimizu KK. 2017. Genome assembly and annotation of Arabidopsis halleri, a model for heavy metal hyperaccumulation and evolutionary ecology. Molecular Ecology Resources 17, 1025-1036.

Buckley J, Holub EB, Koch MA, Vergeer P, Mable BK. 2018. Restriction associated DNA-genotyping at multiple spatial scales in Arabidopsis lyrata reveals signatures of pathogen-mediated selection. BMC Genomics 19, 496.

Bushell C, Spielman M, Scott RJ. 2003. The basis of natural and artificial postzygotic hybridization barriers in Arabidopsis species. The Plant Cell 15, 1430-1442.

Chang PL, Dilkes BP, McMahon M, Comai L, Nuzhdin SV. 2010. Homoeolog-specific retention and use in allotetraploid Arabidopsis suecica depends on parent of origin and network partners. Genome Biology 11,

Clauss M, Koch MA. 2006. Arabidopsis and its poorly known relatives. Trends in Plant Sciences 11, 449–459.

Comai L, Tyagi AP, Winter K, Holmes-Davis R, Reynolds SH, Stevens Y, Byers B. 2000. Phenotypic instability and rapid gene silencing in newly formed arabidopsis allotetraploids. The Plant Cell 12, 1551-1568.

Costa-Nunes P, Pontes O, Preuss SB, Pikaard CS. 2010. Extra views on RNA-dependent DNA methylation and MBD6-dependent heterochromatin formation in nucleolar dominance. Nucleus 1, 254-259.

De La Torre AR, Li Z, Van de Peer Y, Ingvarsson PK. 2017. Contrasting rates of molecular evolution and patterns of selection among gymnosperms and flowering plants. Molecular Biology and Evolution 34, 1363–1377.

Dorofeyev VI. 2002. Cruciferae of European Russia. Turczaninowia 5, 5-114.

Durvasula A, Fulgione A, Gutaker RM, et al. 2017. African genomes illuminate the early history and transition to selfing in Arabidopsis thaliana. Proceedings of the National Academy of Sciences, USA 114, 5213-5218.

Elven R, Murray DF. 2008. New combinations in the Panarctic vascular plant flora. Journal of the Botanical Research Institute of Texas 2, 433-446.

Ernst WHO. 1974. Schwermetallvegetation der Erde. Stuttgart: Fischer.

Exposito-Alonso M, Becker C, Schuenemann VJ, et al. 2018. The rate and potential relevance of new mutations in a colonizing plant lineage. PLoS Genetics 14, e1007155.

Falush D, Stephens M, Pritchard JK. 2007. Inference of population structure using multilocus genotype data: dominant markers and null alleles. Molecular Ecology Notes 7, 574-578.

Forsythe ES, Nelson ADL, Beilstein MA. 2017. Epistatic interactions drive biased gene retention in the face of massive nuclear introgression. BioRxiv 197087, [Preprint.]

Fulgione A, Hancock AM. 2018. Archaic lineages broaden our view on the history of Arabidopsis thaliana, New Phytologist 219, 1194–1198.

Fulgione A, Koornneef M, Roux F, Hermisson J, Hancock A. 2018. Madeiran Arabidopsis thaliana reveals ancient long-range colonization and clarifies demography in Eurasia. Molecular Biology and Evolution 35, 564-574.

1001 Genomes Consortium. 2016. 1,135 genomes reveal the global pattern of polymorphism in Arabidopsis thaliana. Cell 166, 481-491.

German DA, Ebel AL. 2005. Generic placement of Arabidopsis rupicola (Cruciferae). Turczaninowia 8, 5-12.

Gibbard PL. 2002. Quaternary glaciations: extent and chronology. Heidelberg: Elsevier.

Göbel U, Arce AL, He F, Rico A, Schmitz G, de Meaux J. 2018. Robustness of transposable element regulation but no genomic shock observed in interspecific Arabidopsis hybrids. Genome Biology and Evolution **10,** 1403–1415.

Griffin PC. Willi Y. 2014. Evolutionary shifts to self-fertilisation restricted to geographic range margins in North American Arabidopsis lyrata. Ecology Letters 17, 484-490.

Guo X, Li S, Zhang J, et al. 2017. Genome sequencing of 39 Akkermansia muciniphila isolates reveals its population structure, genomic and functional diverisity, and global distribution in mammalian gut microbiotas. BMC Genomics 18, 800.

Guo YL, Zhao X, Lanz C, Weigel D. 2011. Evolution of the S-locus region in Arabidopsis relatives. Plant Physiology 157, 937–947.

Hämälä T, Mattila TM, Leinonen PH, Kuittinen H, Savolainen O. 2017. Role of seed germination in adaptation and reproductive isolation in Arabidopsis lyrata. Molecular Ecology 26, 3484–3496.

Hämäla T, Matila TM, Savolainen O. 2018. Local adaptation and ecological differentiation under selection, migration, and drift in Arabidopsis lyrata. Evolution doi: 10.1111/evo.13502.

Hanikenne M, Talke IN, Haydon MJ, Lanz C, Nolte A, Motte P, Kroymann J, Weigel D, Krämer U. 2008. Evolution of metal hyperaccumulation required cis-regulatory changes and triplication of HMA4. Nature 453, 391-395.

Haudry A, Platts AE, Vello E, et al. 2013. An atlas of over 90,000 conserved noncoding sequences provides insight into crucifer regulatory regions. Nature Genetics 45, 891-898.

He F, Arce AL, Schmitz G, Koornneef M, Novikova P, Beyer A, de Meaux J. 2016. The footprint of polygenic adaptation on stress-responsive cis-regulatory divergence in the Arabidopsis genus. Molecular Biology and Evolution **33**, 2088–2101.

Hedge IC. 1968. Arabidopsis. Flora Iranica 57, 328-334.

Heidel AJ, Ramos-Onsins SE, Wang WK, Chiang TY, Mitchell-Olds T. 2010. Population history in Arabidopsis halleri using multilocus analysis. Molecular Ecology 19, 3364-3379.

Higashi H, Ikeda H, Setoguchi H. 2012. Population fragmentation causes randomly fixed genotypes in populations of Arabidopsis kamchatica in the Japanese Archipelago. Journal of Plant Research 125, 223-233.

Hohmann N. 2011. DNA-barcoding and population genetics of Arabidopsis thaliana's wild relatives. Master thesis, Heidelberg University.

Hohmann N. 2016. Population genomics beyond a model system: the evolutionary history of Arabidopsis thaliana's wild relatives. PhD thesis, Heidelberg University.

Hohmann N, Koch MA. 2017. An Arabidopsis introgression zone studied at high spatio-temporal resolution: interglacial and multiple genetic contact exemplified using whole nuclear and plastid genomes. BMC Genomics 18,

Hohmann N, Schmickl R, Chiang TY, Lučanová M, Kolář F, Marhold K, Koch MA. 2014. Taming the wild: resolving the gene pools of non-model Arabidopsis lineages. BMC Evolutionary Biology 14, 224.

Hohmann N, Wolf EM, Lysak MA, Koch MA. 2015. A time-calibrated road map of Brassicaceae species radiation and evolutionary history. The Plant Cell 27, 2770-2784.

Hu TT, Pattyn P, Bakker EG, et al. 2011. The Arabidopsis lyrata genome sequence and the basis of rapid genome size change. Nature Genetics 43,

Huang CC, Hung KH, Wang WK, Ho CW, Huang CL, Hsu TW, Osada N, Hwang CC, Chiang TY. 2012. Evolutionary rates of commonly used nuclear and organelle markers of Arabidopsis relatives (Brassicaceae). Gene **499.** 194-201.

Huang CH, Sun R, Zeng YH, et al. 2016. Resolution of Brassicaceae phylogeny using nuclear genes uncovers nested radiations and supports convergent morphological evolution. Molecular Biology and Evolution 33, 394-412

Iljinska A, Didukh Y, Burda R, Korotschenko I. 2007. Ecoflora of Ukraine. Kiew: Phytosociocentre Press (in Ukrainian).

Jakobsson M, Hagenblad J, Tavaré S, Säll T, Halldén C, Lind-Halldén C. Nordborg M. 2006. A unique recent origin of the allotetraploid species Arabidopsis suecica: evidence from nuclear DNA markers. Molecular Biology and Evolution 23, 1217-1231.

Jacquemin J, Hohmann N, Buti M, Selvaggi A, Müller T, Koch MA, **Schmid KJ.** 2016. Levels and patterns of genetic diversity differ between two closely related endemic Arabidopsis species. BioRxiv 048785. [Preprint.]

Jordon-Thaden IE, Al-Shehbaz IA, Koch MA. 2013. Species richness of a globally distributed, arctic-alpine genus, Draba L. (Brassicaceae). Alpine Botany 123, 97-106.

Jørgensen MH, Ehrich D, Schmickl R, Koch MA, Brysting AK. 2011. Interspecific and interploidal gene flow in Central European Arabidopsis (Brassicaceae). BMC Evolutionary Biology 11, 346.

Kadota Y. 2007. Arabidopsis umezawana (Brassicaceae), a new species from Mt. Rishirizan, Rishiri Island, Hokkaido, Northern Japan. Japanese Journal of Botany 82, 232-237.

Kagale S, Robinson SJ, Nixon J, et al. 2014. Polyploid evolution of the Brassicaceae during the Cenozoic era. The Plant Cell 26, 2777–2791.

Karl R, Koch MA. 2013. A world-wide perspective on crucifer speciation and evolution: phylogenetics, biogeography and trait evolution in tribe Arabideae. Annals of Botany 112, 983-1001.

Kawagoe T, Kudoh H. 2010. Escape from floral herbivory by early flowering in Arabidopsis halleri subsp. gemmifera. Oecologia 164, 713-720.

Kenta T, Yamada A, Onda Y. 2011. Clinal variation in flowering time and vernalisation requirement across a 3000-M altitudinal range in perennial Arabidopsis kamchatica ssp. kamchatica and annual lowland subspecies kawasakiana. Ecosystem & Ecography S6, 0011.

Kiefer M, Schmickl R, German DA, Mandáková T, Lysak MA, Al-Shehbaz IA, Franzke A, Mummenhoff K, Stamatakis A, Koch MA. 2014. BrassiBase: introduction to a novel knowledge database on Brassicaceae evolution. Plant & Cell Physiology 55, e3.

Koch M, Bishop J, Mitchell-Olds T. 1999. Molecular systematics and evolution of Arabidopsis and Arabis. Plant Biology 1, 529-537.

Koch MA, German DA. 2013. Taxonomy and systematics are key to biological information: Arabidopsis, Eutrema (Thellungiella), Noccaea and Schrenkiella (Brassicaceae) as examples. Frontiers in Plant Science 4, 267.

Koch MA, German DA, Kiefer M, Franzke A. 2018. Database taxonomics as key to modern plant biology. Trends in Plant Science 23, 4-6.

Koch MA, Haubold B, Mitchell-Olds T. 2000. Comparative evolutionary analysis of chalcone synthase and alcohol dehydrogenase loci in Arabidopsis, Arabis, and related genera (Brassicaceae). Molecular Biology and Evolution 17. 1483-1498.

Koch M, Haubold B, Mitchell-Olds T. 2001. Molecular systematics of the Brassicaceae: evidence from coding plastidic matK and nuclear Chs sequences. American Journal of Botany 88, 534-544.

Koch MA, Karl R, German GA. 2016. Underexplored biodiversity of Eastern Mediterranean biota: systematics and evolutionary history of the genus Aubrieta (Brassicaceae). Annals of Botany 119, 39-57.

Koch MA, Matschinger M. 2007. Evolution and genetic differentiation among relatives of Arabidopsis thaliana. Proceedings of the National Academy of Sciences, USA 104, 6272-6277.

Koch MA, Wernisch M, Schmickl R. 2008. Arabidopsis thaliana's wild relatives: an updated overview on systematics, taxonomy and evolution. Taxon 57, 933-943.

Kolár F, Fuxová G, Záveská E, Nagano AJ, Hyklova L, Lucanová M, Kudoh H. Marhold K. 2016a. Northern glacial refugia and altitudinal niche divergence shape genome-wide differentiation in the emerging plant model Arabidopsis arenosa. Molecular Ecology 25, 3929-3949.

Kolár F, Lucanová M, Záveská E, Fuxová G, Mandáková T, Spaniel S, Senko D, Svitok M, Kolník M, Gudzinskas Z, Marhold K. 2016b. Ecological segregation does not drive the intricate parapatric distribution of diploid and tetraploid cytotypes of the Arabidopsis arenosa group (Brassicaceae). Biological Journal of the Linnean Society 119, 673-688.

Kolník M, Marhold K. 2006. Distribution, chromosome numbers and nomenclature concept of Arabidopsis halleri (Brassicaceae) in the Carpathians. Biologia **61,** 41–50.

Koornneef M. Meinke D. 2010. The development of *Arabidopsis* as a model plant. The Plant Journal 61, 909-921.

Laibach F. 1943. Arabidopsis thaliana (L.) Hevnh. Als Obiekt für genetische und entwicklungsphysiologische Untersuchungen. Botanisches Archiv 44,

Leinonen PH, Remington DL, Savolainen O. 2011. Local adaptation, phenotypic differentiation, and hybrid fitness in diverged natural populations of Arabidopsis lyrata. Evolution 65, 90–107.

Llaurens V, Castric V, Austerlitz F, Vekemans X. 2008. High paternal diversity in the self-incompatible herb Arabidopsis halleri despite clonal reproduction and spatially restricted pollen dispersal. Molecular Ecology 17, 1577-1588.

Lynch M. 2010. Evolution of the mutational rate. Trends in Genetics 268, 345-352.

Lysak MA, Cheung K, Kitschke M, Bures P. 2007. Ancestral chromosomal blocks are triplicated in Brassiceae species with varying chromosome number and genome size. Plant Physiology 145, 402-410.

Lysak MA, Koch MA, Beaulieu JM, Meister A, Leitch IJ. 2009. The dynamic ups and downs of genome size evolution in Brassicaceae. Molecular Biology and Evolution 26, 85-98.

Lysak MA, Koch MA, Pecinka A, Schubert I. 2005. Chromosome triplication found across the tribe Brassiceae. Genome Research 15, 516–525.

Mable BK, Hagmann J, Kim ST, Adam A, Kilbride E, Weigel D, Stift M. 2017. What causes mating system shifts in plants? Arabidopsis lyrata as a case study. Heredity 118, 110.

Mable BK, Robertson AV, Dart S, Di Berardo C, Witham L. 2005. Breakdown of self-incompatibility in the perennial Arabidopsis lyrata (Brassicaceae) and its genetic consequences. Evolution **59**, 1437–1448.

Mandákova T. Li Z. Barker MS. Lvsak MA. 2017. Diverse genome organization following 13 independent mesopolyploid events in Brassicaceae contrasts with convergent patterns of gene retention. The Plant Journal 91, 3-21.

Mandákova T, Lysak MA. 2018. Post-polyploid diploidization and diversification through dysploid changes. Current Opinion in Plant Biology

Marhold K, Perný M, Kolník M. 2003. Miscellaneous validations in Cruciferae and Crassulaceae. Willdenowia 33, 69-70.

Mattila TM, Tyrmi J, Pyhäjärvi T, Savolainen O. 2017. Genome-wide analysis of colonization history and concomitant selection in Arabidopsis lyrata. Molecular Biology and Evolution 34, 2665–2677.

Meyer CL, Kostecka AA, Soumitou-Laprade P, Créach A, Castric V, Pauwels M, Frérot H. 2009. Variability of zinc tolerance among and within populations of the pseudometallophyte species Arabidopsis halleri and possible role of directional selection. New Phytologist 185, 130-142.

Meyerowitz EM. 2001. Prehistory and history of Arabidopsis research. Plant Physiology 125, 15-19.

Miller M, Song Q, Shi X, Juenger TE, Chen ZJ. 2015. Natural variation in timing of stress-responsive gene expression predicts heterosis in intraspecific hybrids of Arabidopsis. Nature Communications 6, e7453.

Morgan CM, Zhang H, Bomblies K. 2017. Are the effects of elevated temperature on meiotic recombination and thermotolerance linked via the axis and synaptonemal complex? Proceedings of the Royal Society B: Biological Sciences 372, doi: 10.1098/rstb.2016.0470.

Mummenhoff K, Hurka K. 1995. Allopolyploid origin of Arabidopsis suecica (Fries) Norrlin: evidence from chloroplast and nuclear genome markers. Botanica Acta 108, 449-456.

Nasrallah ME, Liu P, Sherman-Broyles S, Boggs NA, Nasrallah JB. 2004. Natural variation in expression of self-incompatibility in Arabidopsis thaliana: implications for the evolution of selfing. Proceedings of the National Academy of Sciences, USA 101, 16070-16074.

Nasrallah ME, Yogeeswaran K, Snyder S, Nasrallah JB. 2000. Arabidopsis species hybrids in the study of species differences and evolution of amphiploidy in plants. Plant Physiology 124, 1605-1614.

Novikova PY, Hohmann N, Nizhynska V, et al. 2016. Sequencing of the genus Arabidopsis identifies a complex history of nonbifurcating speciation and abundant trans-specific polymorphism. Nature Genetics 48, 1077–1082.

Novikova PY, Hohmann N, Van de Peer Y. 2018. Polyploid Arabidopsis species originated around recent glaciation maxima. Current Opinion in Plant Biology 42, 8-15.

Novikova PY, Tsuchimatsu T, Simon S, et al. 2017. Genome sequencing reveals the origin of the allotetraploid Arabidopsis suecica. Molecular Biology and Evolution 34. 957-968.

O'Kane SL, Al-Shehbaz IA. 1997. A synopsis of Arabidopsis (Brassicaceae). Novon 7. 323-327.

O'Kane SL, Al-Shehbaz IA. 2003. Phylogenetic position and generic limits of Arabidopsis (Brassicaceae) based on sequences of nuclear ribosomal DNA. Annals of the Missouri Botanical Gardens 90, 603-612.

O'Kane SL, Schaal BA, Al-Shehbaz IA. 1996. The origins of Arabidopsis suecica (Brassicaceae) as indicated by nuclear rDNA sequences. Systematic Botany 21, 559-566.

Ossowski S, Schneeberger K, Lucas-Lledó JI, Warthmann N, Clark RM, Shaw RG, Weigel D, Lynch M. 2010. The rate and molecular spectrum of spontaneous mutations in Arabidopsis thaliana. Science 327, 92 - 94

Paape T, Briskine RV, Lischer HEL, et al. 2018. Patterns of polymorphism, selection and linkage disequilibrium I the subgenomes of the allopolyploid Arabidopsis kamchatica. BioRxiv 248195. [Preprint.]

Paape T, Hatakeyama M, Shimizu-Inatsugi R, Cereghetti T, Onda Y, Kenta T, Sese J, Shimizu KK. 2016. Conserved but attenuated parental gene expression in allopolyploids: constitutive zinc hyperaccumulation in the allotetraploid Arabidopsis kamchatica. Molecular Biology and Evolution 33, 2781-2800

Pauwels M, Saumitou-Laprade P, Holl AC, Petit D, Bonnin I. 2005. Multiple origin of metallicolous populations of the pseudometallophyte Arabidopsis halleri (Brassicaceae) in central Europe: the cpDNA testimony. Molecular Ecology 14, 4403-4414.

Pauwels M, Vekemans X, Godé C, Frérot H, Castric V, Saumitou-Laprade P. 2012. Nuclear and chloroplast DNA phylogeography reveals vicariance among European populations of the model species for the study of metal tolerance, Arabidopsis halleri (Brassicaceae). New Phytologist 193,

Pontvianne F, Blevins T, Chandrasekhara C, Feng W, Stroud H, Jacobsen SE, Michaels SD, Pikaard CS. 2012. Histone methyltransferases regulating rRNA gene dose and dosage control in Arabidopsis. Genes & Development 26, 945-957.

Provart NJ, Alonso J, Assmann SM, et al. 2016. 50 years of Arabidopsis research: highlights and future directions. New Phytologist 209, 921-944.

Ross-Ibarra J, Wright SI, Foxe JP, Kawabe A, DeRose-Wilson L, Gos G, Charlesworth D, Gaut BS. 2008. Patterns of polymorphism and demographic history in natural populations of Arabidopsis lyrata. PLoS One

Roux C, Castric V, Pauwels M, Wright SI, Saumitou-Laprade P, Vekemans X. 2011. Does speciation between Arabidopsis halleri and Arabidopsis lyrata coincide with major changes in a molecular target of adaptation? PLoS One 6, e26872.

Säll T, Jakobsson M, Lind-Halldén C, Halldén C. 2003. Chloroplast DNA indicates a single origin of the allotetraploid Arabidopsis suecica. Journal of Evolutionary Biology 16, 1019-1029.

Sato Y, Kudoh H. 2017. Fine-scale frequency differentiation along a herbivory gradient in the trichome dimorphism of a wild Arabidopsis. Ecology and Evolution 7, 2133–2141.

Schmickl R, Jorgenson M, Brysting A, Koch MA. 2008. Phylogeographic implications for North American arctic Arabidopsis lyrata. Plant Ecology & Diversity 1, 245-254.

Schmickl R, Jørgensen MH, Brysting AK, Koch MA. 2010. The evolutionary history of the Arabidopsis lyrata complex: a hybrid in the amphi-Beringian area closes a large distribution gap and builds up a genetic barrier. BMC Evolutionary Biology 10, 98.

Schmickl R. Koch MA. 2011. Arabidopsis hybrid speciation processes. Proceedings of the National Academy of Sciences, USA 108, 14192–14197.

Schmickl R. Paule J. Klein J. Marhold K. Koch MA. 2012. The evolutionary history of the Arabidopsis arenosa complex: diverse tetraploids mask the Western Carpathian center of species and genetic diversity. PLoS One 7. e42691.

Sharbel TF, Haubold B, Mitchell-Olds T. 2000. Genetic isolation by distance in *Arabidopsis thaliana*: biogeography and postglacial colonization of Europe. Molecular Ecology 9, 2109-2118.

Shimizu KK, Fujii SH, Marhold K, Watanabe K, Kudoh H. 2005. Arabidopsis kamchatica (Firsh. ex DC.) K. Shimizu & Kudoh and A. kamchatica subsp. kawasakiana (Makino) K. Shimizu & Kudoh, new combinations. Acta Phytotaxonomica et Geobotanica 156, 163-172.

Shimizu KK. Kudoh H. Kobavashi MJ. 2011. Plant sexual reproduction during climate change: gene function in natura studied by ecological and evolutionary systems biology. Annals of Botany 108, 777-787.

Shimizu KK, Tsuchimatsu T. 2015. Evolution of selfing: recurrent patterns in molecular adaptation. Annual Review of Ecology, Evolution and Systematics 46, 593-622.

Shimizu-Inatsugi R, Lihová J, Iwanaga H, Kudoh H, Marhold K, Savolainen O, Watanabe K, Yakubov VV, Shimizu KK. 2009. The allopolyploid Arabidopsis kamchatica originated from multiple individuals of Arabidopsis lyrata and Arabidopsis halleri. Molecular Ecology 18, 4024–4048.

Solhaug EM, Ihinger J, Jost M, Gamboa V, Marchant B, Bradford D, Doerge RW, Tyagi A, Replogle A, Madlung A. 2016. Environmental regulation of heterosis in the allopolyploid Arabidopsis suecica. Plant Physiology 170, 2251-2263.

Steets JA, Takebayashi N, Byrnes JM, Wolf DE. 2010. Heterogeneous selection on trichome production in Alaskan Arabidopsis kamchatica (Brassicaceae). American Journal of Botany 97, 1098-1108.

Stein RJ, Höreth S, de Melo JR, Syllwasschy L, Lee G, Garbin ML, Clemens S, Krämer U. 2017. Relationships between soil and leaf mineral composition are element-specific, environment-dependent and geographically structured in the emerging model Arabidopsis halleri. New Phytologist 213, 1274-1286.

Takou M, Wieters B, Kopriva S, Coupland G, Linstädter A, De Meaux **J.** 2018. Linking genes with ecological strategies in *Arabidopsis thaliana*. Journal of Experimental Botany (in press).

Tang C, Toomajian C, Sherman-Broyles S, Plagnol V, Guo YL, Hu TT, Clark RM, Nasrallah JB, Weigel D, Nordborg M. 2007. The evolution of selfing in Arabidopsis thaliana. Science 317, 1070-1072.

Tedder A, Hoebe P, Ansell SW, Mable BK. 2010. Using chloroplast tmF pseudogenes for phylogeography in Arabidopsis lyrata. Diversity 2, 653-678.

Tian L. Li X. Ha M. Zhang C. Chen ZJ. 2014. Genetic and epigenetic changes in a genomic region containing MIR172 in *Arabidopsis* allopolyploids and their progenitors. Heredity 112, 207-214.

Tsuchimatsu T, Kaiser P, Yew CL, Bachelier JB, Shimizu KK. 2012. Recent loss of self-incompatibility by degradation of the male component in allotetraploid Arabidopsis kamchatica. PLoS Genetics 8, e1002838.

Turland NJ, Wiersema JH, Barrie FR, et al. 2018. International code of nomenclature for algae, fungi, and plants (Shenzhen code). Regnum Vegetabile 159, XXXVII, 1-254.

Turner TL, Bourne EC, Von Wettberg EJ, Hu TT, Nuzhdin SV. 2010. Population resequencing reveals local adaptation of Arabidopsis lyrata to serpentine soils. Nature Genetics 42, 260-263.

Turner TL, von Wettberg EJ, Nuzhdin SV. 2008. Genomic analysis of differentiation between soil types reveals candidate genes for local adaptation in Arabidopsis lyrata. PLoS One 3, e3183.

Van Rossum F, Bonnin I, Fenart S, Pauwels M, Petit D, Saumitou-**Laprade P.** 2004. Spatial genetic structure within a metallicolous population of Arabidopsis halleri, a clonal, self-incompatible and heavy-metal-tolerant species. Molecular Ecology **13,** 2959–2967.

Veatch-Blohm ME, Roche BM, Dahl EE. 2017. Serpentine populations of Arabidopsis lyrata show evidence for local adaptation in response to nickel exposure at germination and during juvenile growth. Environmental and Experimental Botany 138, 1-9.

Vekemans X. Poux C. Goubet PM. Castric V. 2014. The evolution of selfing from outcrossing ancestors in Brassicaceae: what have we learned from variation at the S-locus? Journal of Evolutionary Biology 27, 1372-1385

Vergeer P, Kunin W. 2011. Life history variation in Arabidopsis lyrata across its range: effects of climate, population size and herbivory. Oikos **120.** 979–990.

Warwick SI, Al-Shehbaz IA, Sauder CI. 2006. Phylogenetic position of Arabis arenicola and generic limits of Aphragmus and Eutrema (Brassicaceae) based on sequences of nuclear ribosomal DNA. Canadian Journal of Botany 84, 269-281.

Wasowicz P, Pauwels M, Pasierbinski A, Przedpelska-Wasowicz EM, Babst-Kostecka AA, Saumitou-Laprade P, Rostanski A. 2016. Phylogeography of Arabidopsis halleri (Brassicaceae) in mountain regions of Central Europe inferred from cpDNA variation and ecological niche modelling. PeerJ 4, e1645.

Weigel D. 2012. Natural variation in Arabidopsis: from molecular genetics to ecological genomics. Plant Physiology 158, 2-22.

Willi Y, Fracassetti M, Zoller S, Van Buskirk J. 2018. Accumulation of mutational load at the edges of a species range. Molecular Biology and Evolution 35. 781-791.

Wos G. Willi Y. 2018. Thermal acclimation in *Arabidopsis lyrata*: genotypic costs and transcriptional changes. Journal of Evolutionary Biology 31, 123-135.

Wright KM, Arnold B, Xue K, Šurinová M, O'Connell J, Bomblies K. 2015. Selection on meiosis genes in diploid and tetraploid Arabidopsis arenosa. Molecular Biology and Evolution 32, 944-955.

Wu Q, Han TS, Chen X, Chen JF, Zou YP, Li ZW, Xu YC, Gu YL. 2017. Long-term balancing selection contributes to adaptation in Arabidopsis and its relatives. Genome Biology 15, 217.

Yant L. Bomblies K. 2017. Genomic studies of adaptive evolution in outcrossing Arabidopsis species. Current Opinion in Plant Biology 36, 9-14.

Yant L. Hollister JD. Wright KM. Arnold BJ. Higgins JD. Franklin FCH, Bomblies K. 2013. Meiotic adaptation to genome duplication in Arabidopsis arenosa. Current Biology 23, 2151-2156.

Zasadni J, Klapyta P. 2014. The Tatra Mountains during the Last Glaciation Maximum. Journal of Maps 10, 440-456.

Zou YP, Hou XH, Wu Q, et al. 2017. Adaptation of Arabidopsis thaliana to the Yangtze River basin. Genome Biology 18, 239.