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# 1 Genetic variation, environment and demography intersect to shape

# 2 Arabidopsis defense metabolite variation across Europe

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# 12 Abstract

Plants face a variety of challenges within their ever-changing environment. Diverse metabolites
are central to the plants ability to overcome these challenges. Understanding the environmental
and genetic factors influencing the variation in specialized metabolites is the key to understand

16 how plants survive and develop under changing environments. Here we measure the variation in

17 specialized metabolites across a population of 797 natural *Arabidopsis thaliana* accessions. We

18 show a combination of geography, environmental parameters, demography, and different genetic

19 processes that creates a specific pattern in their accumulation and distribution. By identifying and

20 tracking causal polymorphisms at multiple loci controlling metabolites variation we show that

- 21 each locus displays extensive allelic heterogeneity with signatures of both parallel and
- 22 convergent evolutionary processes. These loci combine epistatically and show differing

23 relationships to environmental parameters leading to different distributions. This provides a

- 24 detailed perspective about the complexity of the forces and mechanisms that shape the
- accumulation and distribution of a family of specialized metabolites critical for plant fitness.

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- 28

#### 29 Introduction

The biotic and abiotic components of a plant's habitat/environment are continuously changing. 30 This creates a complex system to which a plant must develop adaptation strategies to ensure 31 survival and reproduction. Metabolites are frequent keys to these strategies, involving the 32 33 production and accumulation of different metabolites from signaling hormones, primary 34 metabolites and a wide array of multi-functional specialized metabolites (Erb & Kliebenstein, 2020; Hanower & Brzozowska, 1975; Hayat et al., 2012; Kim et al., 2012; D J Kliebenstein, 35 36 2004; Malcolm, 1994; Thakur & Rai, 1982; Wolters & Jürgens, 2009; Yang, Lin, & Kao, 2000). 37 The complete suite of these metabolites helps to determine the plants survival and development. 38 A complication in the plants ability to create an optimal blend of metabolite-based strategies is the fact that individual specialized metabolites can have contrasting effects in a complex 39 40 environment. For example, individual specialized metabolites can provide defense against some attackers while simultaneously causing sensitivity to other biotic attackers or abiotic stresses 41 42 (Agrawal, 2000; Bialy, Oleszek, Lewis, & Fenwick, 1990; Erb & Kliebenstein, 2020; Futuyma & Agrawal, 2009; Hu et al., 2018; Lankau, 2007; Opitz & Müller, 2009; Uremis, Arslan, 43 Sangun, Uygur, & Isler, 2009; Züst & Agrawal, 2017). This creates offsetting ecological benefits 44 and costs for individual metabolites that when summed across all the metabolites means that 45 46 there are complex selective pressures driving the differentiation of metabolic profiles within a species and shaping genetic variation within and between populations depending on the diverse 47 challenges faced (Fan, Leong, & Last, 2019; R. Kerwin et al., 2015; Malcolm, 1994; Sønderby, 48 Geu-Flores, & Halkier, 2010; Szakiel, Paczkowski, & Henry, 2011; Wentzell & Kliebenstein, 49 2008; Züst et al., 2012). 50

Recent decades have seen significant advances in the identification of the genetic variation creating this metabolic variation. A common theme developing from these studies is that the metabolic variation within and between species is the result of structural variation at the enzymes responsible for the chemical structures (Chan, Rowe, Corwin, Joseph, & Kliebenstein, 2011; Chan, Rowe, & Kliebenstein, 2010; Fan et al., 2019; Kroymann, Donnerhacke, Schnabelrauch, & Mitchell-Olds, 2003; Moore et al., 2019; Schilmiller, Pichersky, & Last, 2012). These structural variants and the resulting chemical variation strongly influence plant fitness in

response to a broad range of biotic interactions including at least herbivores, but also other plant

species and other members of the same plant species (Bednarek & Osbourn, 2009; Brachi et al., 59 2015; R. E. Kerwin et al., 2017; R. Kerwin et al., 2015; Lankau & Kliebenstein, 2009; Lankau & 60 Strauss, 2007, 2008; Lankau, 2007). Most mechanistic studies of natural variation in specialized 61 metabolism have focused on apparent biallelic phenotypic variation linked to loss-of-function 62 variants. However, it is not clear if biallelic genetic causation is true when extended to a large 63 64 collection of individuals from wide-ranging populations within a species. If selective pressures are sufficiently strong and non-linear, it is possible to have repeated and independent generation 65 66 of structural variants creating the same metabolic variation. This raises the possibility for chemical variation within a species to show hallmarks of parallel evolution, wherein 67 phenotypically similar variants independently arise from the same genetic background. Equally it 68 may be possible to find within-species convergent evolution, where different allele with identical 69 70 metabolic consequences arise from independent genetic backgrounds through different mechanisms. Because these genetic processes are occurring simultaneously with neutral 71 72 demographic processes like migration, there is a need to better understand how the intersection 73 of environmental pressure, demography and genomic complexity gives rise to the pattern of 74 metabolic variation across a plant species.

75 To better understand how genomic variation, demography and environmental pressures shape the 76 variation of specialized metabolism within a species, we used the model Glucosinolates (GSLs) 77 pathway. GSLs are a diverse class of specialized metabolites produced in the order Brassicales, including the model plant Arabidopsis (Arabidopsis thaliana), that show extensive variation 78 between and within species across the order (Bakker, Traw, Toomajian, Kreitman, & Bergelson, 79 80 2008; Benderoth et al., 2006; Brachi et al., 2015; Chan et al., 2010; Daxenbichler et al., 1991; Halkier & Gershenzon, 2006; R. Kerwin et al., 2015; D J Kliebenstein, Gershenzon, & Mitchell-81 Olds, 2001; D J Kliebenstein, Kroymann, et al., 2001; D J Kliebenstein, Lambrix, Reichelt, 82 83 Gershenzon, & Mitchell-Olds, 2001; James E. Rodman, Kruckeberg, & Al-Shehbaz, 1981; James Eric Rodman, 1980; Sønderby et al., 2010; Wright, Lauga, & Charlesworth, 2002). GSLs 84 consist of a common core structure with a highly diverse side chain that determines the GSLs 85 biological activity in defense, growth, development and abiotic stress resistance (Beekwilder et 86 al., 2008; Hansen et al., 2008; Hasegawa, Yamada, Kosemura, Yamamura, & Hasegawa, 2000; 87 Katz et al., 2020; Katz, Nisani, Sela, Behar, & Chamovitz, 2015; Malinovsky et al., 2017; 88 89 Salehin et al., 2019; Yamada et al., 2003). The Arabidopsis-GSL system is an optimal model to

study the species wide processes driving specialized metabolite variation because the identity of 90 the whole biosynthetic pathway is known, including the major causal loci for natural variation 91 92 (Benderoth et al., 2006; Brachi et al., 2015; Chan et al., 2011, 2010; Hansen, Kliebenstein, & Halkier, 2007; D J Kliebenstein, Gershenzon, et al., 2001; D. Kliebenstein, Pedersen, Barker, & 93 Mitchell-Olds, 2002; Daniel J Kliebenstein, Figuth, & Mitchell-Olds, 2002; Kroymann & 94 95 Mitchell-Olds, 2005; Pfalz, Vogel, Mitchell-Olds, & Kroymann, 2007; Sønderby et al., 2010; Wentzell et al., 2007). These major loci, have been proven to influence Arabidopsis fitness and 96 97 can be linked to herbivore pressure (Brachi et al., 2015; Hansen et al., 2008; Jander, Cui, Nhan, Pierce, & Ausubel, 2001; R. E. Kerwin et al., 2017; R. Kerwin et al., 2015; Züst et al., 2012). 98 Beyond the major causal loci, there is also evidence from genome wide association studies for 99 highly polygenic variation in the genetic background that further contributes to modulating GSL 100 101 variation (Chan et al., 2011). The public availability of over 1000 widely distributed accessions with genomic sequences provides the ability to phenotype GSL variation across a large spatial 102 103 scale and query the distribution and relationship of causal haplotypes at the major GSL causal loci. 104

105 In Arabidopsis and other Brassicas, the main GSLs are Methionine-derived, Aliphatic, GSLs. Genetic variation in Aliphatic GSLs structure is controlled by natural variation at three loci, GS-106 107 Elong, GS-AOP and GS-OH with these three loci combining to create a dominant Aliphatic GSL 108 chemotype. In addition to these expressed loci, there is a large suite of loci that can modify these dominant patterns (Brachi et al., 2015; Chan et al., 2011, 2010). GS-Elong differentially 109 110 elongates the Methionine side chain by structural variation influencing the expression of divergent methylthioalkylmalate synthase enzymes (MAM) that add carbons to the side chain 111 (Abrahams, Pires, & Schranz, 2020). In Arabidopsis, MAM2 catalyzes the addition of two 112 carbons to the side chain, creating GSLs with 3 carbon side chains. MAM1 catalyzes the addition 113 114 of three carbons to make GSLs with 4 carbon side chains (Figure 1). MAM3 (also known as MAM-L) catalyzes the addition of up to 6 carbons (D J Kliebenstein, Lambrix, et al., 2001; 115 Kroymann et al., 2003; Mithen, Clarke, Lister, & Dean, 1995). The core pathway leads to the 116 creation of the methylthio GSL (MT). Then, the MT will be converted to a methylsulfinyl 117 (MSO) with a matching number of carbons (Giamoustaris & Mithen, 1996; Hansen et al., 2007). 118 Structural variation at the GS-AOP locus leads to differential modification of the MSO by 119 120 differential expression of a family of 2-oxoacid-dependent dioxygenases (20DD). The AOP2





Figure 1: Aliphatic GSL biosynthesis pathway. Short names and structures of the GSLs are in black. Genes encoding the causal enzyme for each reaction (Arrow) are in grey. GS-OX is a gene family of five or more genes. OH-But= 2-OH-3-Butenyl.

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enzyme removes the MSO moiety leaving an alkenyl sidechain, while AOP3 leaves a hydroxyl

moiety. Previous work has suggested three alleles of GS-AOP: AOP3 expressing, AOP2

expressing and a null allele (i.e. Col-0 and similar accessions) with nonfunctional copies of

AOP2 and AOP3 leading to MSO accumulation, the AOP substrate (Figure 1) (Chan et al., 2010;

130 D J Kliebenstein, Kroymann, et al., 2001; D J Kliebenstein, Lambrix, et al., 2001; Mithen et al.,

131 1995). The 4C alkenyl side-chain can be further modified by adding a hydroxyl group at the 2C

via the GS-OH 2-ODD (Figure 1) (Hansen et al., 2008). In spite of the evolutionary distance,

independent variation at the same three loci influence the structural diversity in Aliphatic-GSLs

within Brassica, Streptanthus and Arabidopsis (D J Kliebenstein & Cacho, 2016; Lankau &
Kliebenstein, 2009). For example the C3 MAM in Arabidopsis and Brassica represent two
independent lineages as are the MAMs responsible for C4 GSLs, in fact the MAM locus contains
at least three independent lineages that recreate the same length variation (Abrahams et al.,
2020). This indicates repeated evolution across species, but it is not clear how frequently these
loci are changing within a single species or how ecological or demographic processes may shape
within-species variation at these loci.

141 In this work we described GSL variation in seeds of a collection of 797 *Arabidopsis thaliana* 

142 natural accessions collected from locations across Europe. The amounts of GSLs can vary across

143 different tissues and life stages, but there is a strong correlation in the type of Aliphatic GSL

144 produced across tissues because the major causal effect loci are not plastic due to structural

145 variation (Brown, Tokuhisa, Reichelt, & Gershenzon, 2003; D J Kliebenstein, Gershenzon, et al.,

146 2001; D J Kliebenstein, Kroymann, et al., 2001; Petersen, Chen, Hansen, Olsen, & Halkier,

147 2002). Thus, the seeds chemotype is the same as the leaves. Further, seeds have the highest level

148 of GSLs in Arabidopsis and they are stable at room temperature until germination, hence making

seeds a perfect tissue to survey variation. Further, GSLs are known to be important for seed

defenses against herbivores and pathogens (Raybould & Moyes, 2001). By measuring GSLs in

seeds, we identified that the distribution of GSLs and the causal alleles are influenced by a

diverse set of factors with a primary contribution by geography, environmental parameters and

their interaction. We describe here the complex genetic architecture of the three main causal loci

responsible for the GSL composition, show how it effects the actual phenotype, and how it

evolved. Interestingly, the combination of these elements reveal that several evolutional

mechanisms are involved in shaping the GSL variation and distribution, and the integration ofthem result in the pattern described here.

158

159 **Results** 

#### 160 GSL variation across Europe

To investigate the genetic, environmental and demographic parameters influencing the
distribution of Arabidopsis GSL chemotypes, we measured GSLs from seeds of a collection of

797 Arabidopsis thaliana natural accessions (The 1001 Genomes Consortium, 2016). These 163 Arabidopsis accessions were collected from different geographical locations, mainly in and 164 165 around Europe. 23 different GSLs were detected and quantified identifying a wide diversity in composition and amount among the natural accessions with a median heritability of 83%, 166 ranging from 34% to 93% (Supplemental Table 1). To summarize the GSL variation among the 167 accessions we performed principal component analyses (PCA) on the accumulation of all the 168 individual GSLs across the accessions as an unbiased first step. The first two PCs only captured 169 33% of the total variation with PC1 describing GSLs with 4 and 7 carbons and PC2 mainly 170 capturing GSLs with 8 carbons in their side chain (supp. Figure 1). Previous work using a 171 collection of predominantly central European accessions had suggested a simple continental 172 gradient chain-elongation variation from the south-west to the north-east (Brachi et al., 2015; 173 174 Züst et al., 2012). To assess if this was still apparent in this larger collection, we plotted the accessions based on their geographical locations, and colored them based on their PC1 and PC2 175 176 scores that are linked to chain elongation variation (Figure 2A and supp. Figure 2A, respectively). This larger collection shows that there is not a single gradient shaping GSL 177 178 diversity across Europe (Figure 2A). Instead the extended sampling of accessions around the 179 Mediterranean in this collection shows that the SW to NE pattern reiterates within the Iberian 180 Peninsula.



#### 181

Figure 2: **GSL variation across Europe is dominated by two loci.** A. The accessions are plotted on the map based on their collection site, and colored based on their PC1 score. B.

plotted on the map based on their collection site, and colored based on their PC1 score. B.
 Manhattan plot of GWAS analyses using PC1. Horizontal lines represent 5% significance

185 thresholds using Bonferroni (red) and permutations (blue).

To test which of the major causal loci are detectable in this collection and to identify new 186 187 genomic regions that are associated with the observed GSL variation, we performed genome 188 wide association (GWA, with EMMAX algorithms) analyses using the PC1 and PC2 values. 189 This collection of natural accessions presents a dense variant map and is 3x larger than previous GSL GWA mapping populations. In spite of the large population size, both PC1 and PC2 based 190 191 analyses identified the same two major peaks covering two of the known causal genes controlling GSL diversity (Figure 2B for PC1 GWA analyses, supp. Figure 2B for PC2 GWA 192 193 analyses) (Brachi et al., 2015; Chan et al., 2011, 2010). The largest peak in both cases, is the GS-Elong locus on chromosome 5, containing the MAM1 (AT5G23010), MAM2 and MAM3 194 (AT5G23020) genes. The peak on chromosome 4 is the GS-AOP locus containing the AOP2 and 195 AOP3 genes (AT4G03060 and AT4G03050, respectively). Previous F2, QTL and molecular 196 197 experiments have shown that the genes within GS-AOP and GS-Elong loci are the causal genes for GSL variation within these regions (Benderoth et al., 2006; Brachi et al., 2015; Chan et al., 198 199 2011, 2010; D J Kliebenstein, Gershenzon, et al., 2001; D. Kliebenstein et al., 2002; Daniel J Kliebenstein et al., 2002; Kroymann & Mitchell-Olds, 2005; Pfalz et al., 2007; Wentzell et al., 200 201 2007). Surprisingly, none of the 8 other known natural variants within the GSL biosynthetic 202 pathway were identified by GWA including three that were found with 96 accessions and three 203 that were found with 595 accessions using PC1 and 2 (Brachi et al., 2015; Chan et al., 2011, 204 2010; Daniel J. Kliebenstein, 2009). It is possible that the extended sampling of accessions may 205 have created genomic and demographic issues that influenced this high false-negative error rate 206 where  $\sim 80\%$  of validated natural variants found using multiple RIL populations were missed.

207

#### 208 Complex GSL Chemotypic Variation

One potential complicating factor is that GSL chemotypic variation is best described as a discrete
multimodal distribution involving the epistatic interaction of multiple genes which PCA's linear
decomposition cannot accurately capture (Figure 1). To test if PCA was inaccurately describing
GSL chemotypic variation, we directly called the specific GSL chemotypes in each accession.
Using Arabidopsis QTL mapping populations and GWA, we have shown that the GS-AOP,
Elong and OH loci determine seven discrete chemotypes, 3MSO, 4MSO, 3OHP, 4OHB, Allyl,
3-Butenyl, 2-OH-3-Butenyl, that can be readily assigned from GSLs phenotypic data (Brachi et

al., 2015; Chan et al., 2011, 2010; D J Kliebenstein, Gershenzon, et al., 2001). Using accessions 216 with previously known chemotypes and genotypes, we developed a phenotypic classification 217 scheme to assign the chemotype for each accession (Figure 3, for details see methods and supp. 218 Figures 3-5, for structures see Figure 1 and supp. Table 1). Since the Aliphatic GSLs 219 composition in the seeds reliably indicate the GSL structural composition in the other plant's life 220 221 stages and tissues, assigning a chemotype for each accession based on the seeds composition is expected to be highly stable across tissues of the same accession (Brown et al., 2003; Chan et al., 222 2011, 2010; D J Kliebenstein, Gershenzon, et al., 2001; D J Kliebenstein, Kroymann, et al., 223 2001). Most accessions were classified as 2-OH-3-Butenyl (27%) or Allyl (47%) with lower 224 frequencies for the other chemotypes. Mapping the chemotypes on Europe showed that the PCA 225 decomposition was missing substantial information on GSL chemotype variation (Figure 3). 226 227 Instead of a continuous distribution across Europe, the chemotype classifications revealed specific geographic patterns. Central and parts of northern Europe were characterized by a high 228 229 variability involving the co-occurrence of individuals from all chemotypes. In contrast, southern Europe, including the Iberian Peninsula, Italy and the Balkan, has two predominant chemotypes, 230 231 Allyl or 2-OH-3-Butenyl, that are separated by a sharp geographic partitioning (Figure 3, and supp. Figure 6). The few accessions in southern Europe belonging to other chemotypes were all 232 233 accessions previously identified as having genomes identical to accessions in central Europe, suggesting that they are likely stock center seed contaminations (The 1001 Genomes 234 235 Consortium, 2016). Uniquely, Swedish accessions displayed a striking presence of almost solely Allyl chemotypes that was not mirrored on the eastern coast of the Baltic Sea (Finnish, 236 237 Lithuanian, Latvian or Estonian accessions). Directly assigning GSL variation by discrete chemotypes provided a more detailed image not revealed by PCA decomposition. Further, the 238 239 different chemotypic to geographic patterns suggests that there may be different pressures 240 shaping GSL variation particularly when comparing central and southern Europe.

Figure 3: Phenotypic classification based on GSL content. A. Using the GSL accumulation, 241 242 each accession was classified to one of seven aliphatic short chained GSL chemotypes based on the enzyme functions as follows: MAM2, AOP null: classified as 3MSO dominant, colored in 243 yellow. MAM1, AOP null: classified as 4MSO dominant, colored in pink. MAM2, AOP3: 244 245 classified as 3OHP dominant, colored in green. MAM1, AOP3: classified as 4OHB dominant, colored in light blue. MAM2, AOP2: classified as Allyl dominant, colored in blue. MAM1, 246 AOP2, GS-OH non-functional: classified as 3-Butenyl dominant, colored in black. MAM1, 247 AOP2, GS-OH functional: classified as 2-OH-3-Butenyl dominant, colored in red. The 248

- 249 accessions were plotted on a map based on their collection sites and colored based on their
- 250 dominant chemotype. B. The coloring scheme with functional GSL enzymes in the aliphatic
- 251 GSL pathway is shown with the percentage of accessions in each chemotypes (out of the total
- 252 797 accessions) shown in each box.



#### 254 Geography and environmental parameters affect GSL variation

Because GSL chemotypes may be more reflective of local environment, we proceed to test if 255 they are associated with weather parameters and landscape conditions. Further, given the 256 difference in chemotype occurrence in central and southern Europe we hypothesized that these 257 258 environmental connections may change between central and southern Europe. For these tests, we 259 chose environmental parameters that capture a majority of the environmental variance and by that may describe the type of ecosystem (Ferrero-Serrano & Assmann, 2019). We assigned each 260 accession the environmental value based on its location. These environmental parameters include 261 geographic proximity (distance to the coast), precipitation descriptors (precipitation of wettest 262 263 and driest month) and temperature descriptors (maximal temperature of warmest month and 264 minimal temperature of coldest month) capture major abiotic pressures as well as provide 265 information about the type of ecosystem in which each accession exists. We ran a multivariate analysis of variance (MANOVA) for each geographic area separately (north and 266 267 central vs south, as shown in supp. Figure 6). This showed significant difference in how the GSL chemotypes associated to the environmental parameters across Europe. This was best illustrated 268 269 by the two dominant chemotypes, Allyl and 2-OH-3-Butenyl, showing opposing relationships to 270 the precipitation in the driest month. In Northern and Central Europe, the Allyl chemotype is 271 more associated with lower precipitation in the driest month, while accessions with 2-OH-3-272 Butenyl as the dominant chemotype are associated with higher precipitation in the driest month. In Southern European accessions, this association is inverted (Figure 4A,B). This suggests that 273 274 the relationship of GSL chemotype to environmental parameters vary across geographic regions 275 of Europe rather than fitting a simple linear model.

As the two main chemotypes in the collection differ by the length of the carbon chain (C3 for 276 277 Allyl, C4 for 2-OH-3-Butenyl), we created a linear model to further check the interaction 278 between each environmental condition to geography in respect to the carbon chain length. Most of the environmental parameters significantly interacted with geography, meaning that the 279 280 relationship of environment to GSL alleles change across geographic areas (supp. Figure 7, for details on the models see methods). Conducting this analysis for each of the geographic areas 281 separately highlighted this by showing that these parameters have different effects on the carbon 282 283 chain length in each of the areas (supp. Figure 7). This was true when the model was run with or

# without ancestral population state being included in the model (The 1001 Genomes Consortium,







Figure 4: Environmental conditions differentially associate with GSLs across geographic 287 location. A. B. The association of the two major chemotypes allyl and 2-OH-3-Butenyl to 288 precipitation values of the driest month. Significance was tested by t-Test, P = 0.00000258 for 289 the North (Slope= 0.01), P = 0.0005521 for the South (Slope= -0.007). C. MANOVA was 290 291 performed for the south and north as indicated in methods section, followed by pairwise comparisons of least-squares means. Numbers indicate chemotypes with significant differences 292 in the North and letters indicate chemotypes with significant differences in the south. OH-But= 293 294 2-OH-3-Butenyl.

295

296 Using a random forest machine learning approach provided similar results with different

environment to chemotype relationships in the north and south (sup. Figure 8), supporting the

298 hypothesis that the GSL chemotype to environment relationships change across regions within

Europe.

300

# 301 The genetic architecture of GSL variation

302 The presence of different GSL chemotype to environmental relationships across Europe raises

the question of how these chemotypes are generated. Are these chemotypes from locally derived

alleles or obtained by the intermixing of widely distributed causal alleles. Further, if there are

multiple alleles, do they display within species convergent or parallel signatures. We focus on

the GS-AOP, GS-Elong, and GS-OH loci, the causal genes creating Arabidopsis GSL

307 chemotypes, and use the available genomic sequences in all of these accessions to investigate the
308 allelic variation in these genes to map the allelic distribution and test the potential for convergent
309 and/or parallel evolution within each locus.

310 GS-Elong: Because the variation in the GS-Elong locus is caused by complex structural variation

in MAM1 and MAM2 that is not resolvable using the available data from short-read genomic

sequence, we used the MAM3 sequence within this locus to ascertain the genomic relationship of

accessions at the causal GS-Elong locus (Kroymann et al., 2003). We aligned the MAM3

sequence from each of the accessions, rooted the tree with the *Arabidopsis lyrata* orthologue

315 (MAMc), and colored the tree tips based on the accessions dominant chemotype.

316 The accessions were distributed across eight distinctive clades with each clade clustering

accessions having either a C3 or C4 status (Figure 5A). The clades C3/C4 status altered across

the tree with three of the clades C3 dominant (MAM2 expressed), and five clades being C4

dominant (MAM1 expressed). Further supporting the use of MAM3 is that the accession

320 assignments to these clades agree with available bacterial artificial chromosome-based

321 sequencing of the GS-Elong region from 15 accessions (Figure 5B). While there are multiple

functional alleles for both C3 and C4 chemotypes, the genomic sequence and phylogeny does not

appear consistent with a simple parallel evolution model where one allele/population is the basis

324 for the independent derivation of all alternative alleles. This is illustrated by the difference in the

genomic arrangement of Clade 4 and 5 which both create C4 GSLs. Clade 4 has a copy of

326 MAM2 and MAM1 while Clade 5 has two copies of MAM1 (Figure 5). It appears that Clade 4 is

the basis for two independent C3 alleles via separate deletions of MAM1 (Clades 1 and 3) and a

separate C4 allele via a deletion of MAM2 (Clade 2, Figure 5). Unfortunately, no long-read

329 sequencing is available in accessions from Clade 6 or 7 and locus-specific de novo alignment of

short-read sequences in these accessions was not able to resolve the regions complexity. Filling

in these clades would be necessary to better understand convergent/parallel events giving rise toGSL chemotypes.

Interestingly, the most basal clade has no copy of MAM2, raising the question of where MAM2 arose (Figure 5). This suggests that true ancestral state(s) of this locus is not represented in this collection and would need to be searched for in other populations of *Arabidopsis thaliana*, if it exists in extant populations.



Figure 5: **MAM3 phylogeny.** A. MAM3 phylogeny of *Arabdopsisi thaliana* accessions, rooted by *Arabidopsis lyrata* MAMc, that is not shown because of distance. Tree tips are colored based on the accession chemotype. The named accessions indicate that GSL-Elong region of these accessions was previously sequenced (Kroymann et. al. 2003). B. The genomic structure of the GSL-Elong regions in the previously sequenced accessions is shown based on Kroymann et. al. 2003. The accession names are colored based on their clades. The color of the name of the accession indicates the clade it belongs. Bright grey arrows represents MAM1 sequences, dashed

arrows represents MAM2 sequences. Dark grey arrows represent MAM3 sequences. The number
to the right of the genomic cartoon represents the number of carbons in the side chain. C.
Collection sites of the accessions, colored by their clade classification (from section A). D. Clade
2 reflection on the map.

349

Using this phylogeny, we investigated the presence of the different GS-Elong haplotypes across 350 351 Europe to ask if each region has a specific allele/clade or if the alleles are distributed across the continent. Specifically, we were interested if the strong C3/C4 partitioning in southern Europe 352 was driven by the creation of local alleles or if this partitioning might contain a wide range of 353 alleles. If the latter is true, this would argue for a selective pressure shaping this C3/C4 divide. 354 We plotted the accessions on the map and colored them based on their GS-Elong clade (Figure 355 356 5C). This showed that the strong  $C_3/C_4$  partition in the Iberian Peninsula contains haplotypes 357 from all the GS-Elong clades except Clade 3 and is not caused by local alleles. This suggests that the strong geographic partitioning of the C3/C4 chemotypes in Iberia may be driven by selective 358 359 pressure causing the partitioning of the chemotypes rather than neutral demographic processes.

360 Shifting focus to all of Europe showed that while most clades were widely distributed across Europe there were a couple over-arching patterns (Figure 5C, and supp. Figure 9). GS-Elong 361 362 clades 1 and 6 follow a pattern that fits with alleles located within the Iberian glacial refugia that then moved north. In contrast the absence of clade 3 from Iberia is more parsimonious with a 363 364 glacial refugia in the Balkans followed by a northward movement wherein it mixed with the other clades. Other clades never moved north and are exclusive to the south as shown by clades 5 365 366 and 7. While these are both C4 clades, other C4 clades like clades 2 and 8 were able to move 367 north (Figure 5D, and supp. Figure 9, respectively). This suggests that there are either differences in their GSL chemotype influencing their distribution or there are neighboring genes known to be 368 under selection in Arabidopsis like FLC (AT5G10140) that may have influenced their 369 370 distribution. In combination, this suggests that a complex demography is involved in shaping the chemotypes identity with some regions, Iberia, showing evidence of local selection while other 371 372 regions, central Europe, possibly showing a blend requiring further work to delineate (supp. Figure 9). 373

GS-AOP: Side chain modification of the core MSO GSL is determined by the GS-AOP locus.Most of the accessions contain a copy of AOP2 and a copy of AOP3, but only one of them will

be functionally expressed (Chan et al., 2010), while in some cases both will be nonfunctional. To
better understand the demography and evolution of the GS-AOP locus, we separately aligned the
AOP2 and AOP3 sequences, rooted each tree with the *Arabidopsis lyrata* orthologue, and
colored the trees tips based on the accessions dominant chemotype.

380 The phylogenetic trees shared a very similar topology, yielding a clear separation between 381 alkenyl (AOP2 expressed) and hydroxyalkyl (AOP3 expressed) accessions. Alkenyl expressing accessions like Cvi-0 with an expressed copy of the AOP2 enzyme formed a single contiguous 382 cluster (Figure 6A). In contrast, hydroxyalkyl accessions clustered into two separate groups with 383 one group of 3OHP dominant accessions partitioning from the rest of the accessions at the most 384 385 basal split in the tree (supp. Figure 10). This haplotype is marked by having an inversion swapping the AOP2 and AOP3 promoters as shown in bacterial artificial chromosome 386 387 sequencing of the Ler-0 accession (Figure 6D) (Chan et al., 2010). The tree also identified a second group of 3OHP dominant accessions located among the alkenyl accessions. Analyzing 388 389 the sequences of these accessions reveals that this small group of 3OHP accessions have a complete deletion of AOP2 and contain only AOP3 (Figure 6E). Thus, there are at least two 390 391 independent transitions from Alkenyl to Hydroxyalkyl GSLs within Arabidopsis, neither of which are related to the Alkenyl to Hydroxyalkyl conversion within Arabidopsis lyrata. 392

The null accessions (MSO dominant chemotypes) were identifiable in all the major clades on the 393 394 tree (supp. Figure 10, middle column of heatmap) suggesting that there are independent LOF 395 mutations that abolish either AOP2 or AOP3. Deeper examination of the sequences of these 396 accessions identified three convergent LOF alleles leading to the MSO chemotype. Most of the null accessions harbor a 5 bps deletion in their AOP2 sequence, that causes a frameshift 397 mutation. This mutation arose within the Alkenyl haplotype and was first reported in the Col-0 398 399 reference genome (Figure 6B) (D J Kliebenstein, Lambrix, et al., 2001). In addition, there are 400 additional independent LOF events arising in both the alkenyl haplotype (e.g. Sp-0, Figure 6C), and within the Ler-0 inversion haplotype (e.g. Fr-2, Figure 6F). Thus, GS-AOP has repeated 401 402 LOF alleles arising within all the major AOP haplotypes suggesting convergent evolution of the MSO chemotype out of both the Alkenyl and Hydroxyalkyl chemotypes. 403

Using the combined chemotype/genotype assignments at GS-AOP, we investigated thedistribution of the alleles across Europe. The Alkenyl haplotype is spread across the entire

- 406 continent. In contrast, the hydroxyalkyl haplotypes are more local. The Ler-like 3OHP haplotype
- 407 is present in only central and north Europe (Figure 6D), while the other 3OHP haplotype,
- 408 possessing only AOP3, is limited to Azerbaijan, along the Caspian Sea (Figure 6E). In contrast to
- 409 the distinct hydroxyalkyl locations, the distribution of the independent LOF null haplotypes
- 410 overlaps with all of them being located within central and north Europe (Figure 6B, C and F).
- 411 The fact that these independently derived LOF alleles are all contiguous suggests that there may
- 412 be a benefit to these alleles specific to Central Europe.





Figure 6: AOP Genomic structure. The genomic structure and causality of the major

- 415 AOP2/AOP3 haplotypes are illustrated. Pink arrows show the AOP2 gene while yellow arrows
- represent AOP3. The black arrows represent the direction of transcription from the AOP2
- 417 promoter as defined in the Col-0 reference genome. Its position does not change in any of the
- regions. The black lines in Sp-0 and Fr-2 presence the position of independent variants creating

premature stop codons. The GSL chemotype for each haplotype is listed to the right with the
 number of the accessions in brackets. The maps show the geographic distribution of the
 accessions from each structure.

422

423 GS-OH: The final major determinant of natural variation in Arabidopsis GSL chemotype is the GS-OH enzyme that adds a hydroxyl group to the 2 carbon on 3-butentyl GSL to create 2-OH-3-424 425 butenyl GSL. Previous work had suggested two GS-OH alleles measurable in the seed, a functional allele in almost all accessions and a non-functional allele caused by active site 426 mutations represented by the Cvi-0 accession (Hansen et al., 2008). Because of functional 427 epistasis, we can only obtain functional phenotypic information from accessions that accumulate 428 429 the GS-OH substrate, 3-butenyl GLS. This identified 11 accessions with a non-functional GS-430 OH. Surveying these 11 accessions in the polymorph database identified multiple independent LOF events. One of these 11 accessions have the Cvi active site mutations, two accessions have 431 432 a shared nonsense SNP that introduce premature stop codons, and two accessions have a complete loss of this gene (Table 1). We could not identify the causal LOF allele in the other six 433 accessions due to sequence quality in the databases. All of these independent GS-OH LOF 434 435 alleles are phylogenetically positioned within groups of accessions that largely do not accumulate 3-butenyl GLS, e.g. 3 carbon or non-alkenyl accessions suggesting that the 436 functional epistasis may be influencing the generation of these alleles. Thus, we searched the 437 438 accessions that do not accumulate 3-butenyl GLS and have effectively hidden the GS-OH 439 function for these GS-OH LOF events (Supp table 2). In each case, the LOF allele is more 440 frequent in the non 4 carbon-alkenyl accessions than expected by random chance. This suggests that there is a bias against 3-butenyl GSL synthesis as the LOF alleles are more frequent when 441 the GS-OH gene is hidden by functional epistasis. This agrees with the fact that the 3-butenyl 442 443 chemotype is the most sensitive to generalist lepidopteran herbivory (Hansen et al., 2008). Thus, 444 these mutations may represent ongoing pseudogenization of the GS-OH gene when it is functionally hidden by epistasis at the GS-AOP and GS-Elong loci. These LOF events would 445 446 then only be displayed upon rare admixture with 2-OH-3-Butenyl accessions.

447

Accession	Type of mutation	Allele structure
Sorbo, Pien	polymorphism at SNP10831302	
Cvi-0	Active site mutation	
IP-Mot-0, IP-Tri-0	Gene deletion	
T670	Independent mutation	
FlyA 3	Independent mutation	
Ting-1	Independent mutation	
T880	Independent mutation	
T710	Independent mutation	
T850	Independent mutation	

449

Table 1: GS-OH structure. The structures of GS-OH in the 3-Butenyl accessions are illustrated.
 These mutations create premature stop codons.

452

#### 453 Discussion

454 Understanding the genetic, demographic and environmental factors that shape variation within a trait in a population is key to understanding trait evolution. In this work we used Aliphatic GSLs 455 in seeds of Arabidopsis thaliana to query how genetics, geography, environment and 456 457 demography intersect to shape chemotypic variation across Europe. We found that 458 environmental conditions, together with geography affect the presence and distribution of 459 chemotypes within the accessions. This was demonstrated by specific traits that were associated 460 with specific environmental conditions, and this association was shifted across the continent. Comparing the associations of traits to specific environmental conditions in central Europe 461 462 versus the south revealed different, sometimes even inverse, behaviors. For example, In the Iberian Peninsula, 2-OH-3-Butenyl was positively associated with potential drought while in 463 Central Europe, it was the opposite GS-Elong allele showing association. This showed that 464

chemotypic variation across Europe is created by a blend of all these processes that differ at the
individual loci and required the simultaneous analysis of genotype and phenotype to fully
interpret.

In contrast to the bimodal distribution in the Aliphatic GSL traits, each of the three major 468 469 Aliphatic GSL loci showed allelic heterogeneity with multiple independent structural variants 470 that recreated the same phenotypic variation. The GS-AOP locus had numerous events with the AOP3 variant of GS-AOP arising via at least two independent events and the Null allele being 471 472 generated at least 15 independent times. The GS-AOP null alleles convergently arose from all the different functional haplotypes. It is less clear if the independent GS-AOP AOP3 alleles 473 474 should be classified as convergent or parallel due to a lack of clarity in what is the ancestral state. 475 Similar to the independent GS-AOP null alleles, there were numerous independent GS-OH LOF 476 variants with at least 9 independent events. While these are parallel GS-OH LOF events because they came from a single functional GS-OH group, their ability to accumulate depends on the 477 478 epistatic silencing of GS-OH by the GS-AOP and GS-Elong loci. The GS-Elong locus also had an extensive level of allelic heterogeneity hallmarked by a shifting expression of the MAM1 or 479 480 MAM2 gene, again with hallmarks of both parallel and convergent processes. Interestingly, at 481 both the GS-AOP and GS-Elong loci, one gain-of-function event (e.g. AOP3 in GS-AOP locus, 482 and MAM2 in GS-Elong locus) is concurrently linked to a loss-of-function of the other gene at 483 the locus (AOP2 and MAM1, respectively). These structural variants are shaped such that the chemotypes show distinct separations without any intermediate phenotypes. This allelic 484 485 heterogeneity is in contrast to previous work on other biotic interactions genes like pathogen 486 resistance gene-for-gene loci that typically have two moderate frequency stable alleles creating 487 the phenotypic variation within the species (Atwell et al., 2010; Corrion & Day, 2001; MacQueen, Sun, & Bergelson, 2016). In other cases alleles of genes involved in biotic defense 488 489 can present more complex patterns, e.g. natural variation in the immune gene ACCELERATED CELL DEATH 6 (ACD6) is caused by a rare allele causing an extreme lesion phenotype. It is not 490 491 yet clear what selective pressures influence ACD6 genetic variation (Todesco et al., 2010; Zhu et al., 2018). The contrast where Aliphatic GSL loci have high levels of allelic heterogeneity for 492 independent and recurrent LOF and GOF events while other resistance genes have more stable 493 494 biallelic variation suggests that there are different selective regimes influencing these loci.

495 Further work is needed to assess the range of allelic heterogeneity in loci controlling resistance496 to diverse biotic traits within the environment.

497

The allelic heterogeneity at these loci illustrates the benefit of simultaneously tracking the 498 phenotype and genotype when working to understand the distribution of trait variation. For 499 500 example, the Iberian Peninsula and the Mediterranean had low variability in Aliphatic GSL chemotype with the chemotypes not overlapping while central/north Europe had high Aliphatic 501 502 GSL diversity with the chemotypes overlapping. At first glance, this contrasts with previous work showing that the Iberian Peninsula and the Mediterranean are more genetically diverse. 503 However, this discrepancy was caused by one of the causal loci. Specifically, the GS-AOP locus 504 is largely fixed as the Alkenyl allele in Iberia/Mediterranean with the alternative GS-AOP alleles 505 506 enriched in central Europe. In contrast to GS-AOP, Iberia and the Mediterranean were highly genetically diverse for the GS-Elong locus and appear to contain all the variation in GS-Elong 507 508 found throughout Europe (The 1001 Genomes Consortium, 2016). Thus, the chemotypic divergence from genomic variation expectations was driven by just the GS-AOP locus. This 509 indicates that the high level of chemotypic variation in central Europe is a blend of alleles that 510 moved from the south (GS-Elong) and alleles that possibly arose locally (GS-AOP, both nulls 511 512 and AOP3). Further, the chemotypes found in any one region appear to be created by a 513 combination of alleles moving across the continent, local generation of new polymorphisms and 514 local selective pressures that shape the chemotypes distribution across the landscape.

515

516 One difficulty in interpreting the evolutionary processes, e.g. parallel v convergent, especially for 517 structural variants illustrated by all the three loci is the complication in properly identifying the 518 ancestral state of the population. While this could typically be done by relying on shared loci 519 with sister species, this is not possible in this case as Arabidopsis lyrata and halleri have genetic variation at GS-Elong and GS-AOP creating the exact same phenotypes. Further, neither of these 520 521 sister species have yet been found to have a functional GS-OH (Heidel, Clauss, Kroymann, 522 Savolainen, & Mitchell-Olds, 2006; Ramos-Onsins, Stranger, Mitchell-Olds, & Aguadé, 2004; 523 Windsor et al., 2005). The MSO chemotypes could be viewed as convergent evolution within a species, as the MSO phenotype independently re-occurred multiple times in the AOP2 and AOP3 524 genetic backgrounds. However, it is not clear how to classify the different AOP3 (AZE v Ler) 525

types as it not clear if the AOP2 or AOP3/Ler haplotype is ancestral within Arabidopsis thaliana. 526 Another option to calling ancestral state is deep sampling in the species but even with these 527 528 accessions, we do not appear to have reached the necessary threshold. For example previous 529 work at the GS-Elong locus had suggested that the Sorbo accession, collected from Tajikistan, was the most likely ancestral state as it had a copy of both MAM1 and MAM2 (Kroymann et al., 530 531 2003). However, the phylogeny with this larger collection of accessions suggested that Sorbo is not ancestral. Further, a recent phylogeny of MAM genes across the Brassicales suggests that 532 MAM2 is an *Arabidopsis thaliana* specific gene with an undefined origin (Abrahams et al., 533 2020). This suggests that to get a better understanding of the ancestral state to define 534 evolutionary processes, especially for loci with allelic heterogeneity and structural variants, we 535 need to broaden our phylogenetic context by deeper sampling within and between species. 536

537

Another complication caused by the allelic heterogeneity and differential selective pressures 538 539 displayed within this system is that we were unable to detect a number of known and validated natural variants that are causal within this population. Specifically, the GWAS with this 540 541 collection of 797 accessions was unable to find 80% of the known causal loci including one of 542 the three major effect loci, GS-OH. Maximizing the number of genotypes and the SNP marker 543 density was unable to overcome the complications imposed by the complex pressures shaping 544 the distribution of these traits. In this system, the optimal path to identifying the causal polymorphisms has instead been a small number of Recombinant Inbred Line populations 545 546 derived from randomly chosen parents. In complex adaptive systems, the optimal solution to 547 identifying causal variants is likely a blend of structured mapping populations and then translating the causal genes from this system to the GWAS results and tracking the causal loci 548 549 directly.

In this work we combined different approaches to uncover some of the parameters shaping the
Aliphatic GSL content across Europe. Widening the size of the population will enable us to
deepen our understanding on the evolutionary mechanisms shaping a phenotype in a population.

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555

#### 556 Methods

#### 557 Plant materiel:

558 Seeds for 1135 Arabidopsis (Arabidopsis thaliana) genotypes were obtained from the 1001

559 genomes catalog of Arabidopsis thaliana genetic variation (https://1001genomes.org/). All

560 Arabidopsis genotypes were grown at 22°C/24°C (day/night) under long-day conditions (16-h of

561 light/8-h of dark). Two independent experiments were performed, each of them included the full

set of genotypes. In the analyses only accessions from Europe and around Europe were included

- 563 (Figure 2A), resulting in an analysis of 797 accessions. List of the accessions can be found in
- supp. Table 1.

#### 565 GSL extractions and analyses:

566 GSLs were measured as previously described (D J Kliebenstein, Gershenzon, et al., 2001; D J Kliebenstein, Kroymann, et al., 2001; D J Kliebenstein, Lambrix, et al., 2001). Briefly, ~3mg of 567 seeds were harvested in 200 µL of 90% methanol. Samples were homogenized for 3 min in a 568 paint shaker, centrifuged, and the supernatants were transferred to a 96-well filter plate with 569 DEAE sephadex. The filter plate with DEAE sephadex was washed with water, 90% methanol, 570 and water again. The sephadex bound GSLs were eluted after an overnight incubation with 571 572 110µL of sulfatase. Individual desulfo-GSLs within each sample were separated and detected by HPLC-DAD, identified, quantified by comparison to standard curves from purified compounds, 573 574 and further normalized to the weight. List of GSLs and their structure are in supplementary table 1. Row GSLs data are in supplementary table 1B. 575

# 576 Statistics, heritability, and data visualization:

Statistical analyses were conducted using R software (https://www.R-project.org/) with the 577 RStudio interface (http://www.rstudio.com/). For each independent GLS, a linear model 578 579 followed by ANOVA was utilized to analyze the effect of accession, replicate, and location in the experiment plate upon the measured GLS amount. Broad-sense heritability (supplementary 580 581 table 1C) for the different metabolites was estimated from this model by taking the variance due to accession and dividing it by the total variance. Estimated marginal means (emmeans) for each 582 583 accession were calculated for each metabolite from the same model using the package emmeans ("CRAN - Package emmeans," n.d.) (supplementary table 1D). Principal component analyses 584

- were done with FactoMineR and factoextra packages (Abdi & Williams, 2010). Data analyses
- and visualization was done using R software with tidyverse (Wickham et al., 2019) and ggplot2
- 587 (Kahle & Wickham, 2013) packages.
- 588 Principal component analyses were done with FactoMineR and factoextra packages (Abdi &
- 589 Williams, 2010).
- 590 Maps were generated using ggmap package ("https://journal.r-project.org/archive/2013-1/kahle-
- 591 wickham.pdf," n.d.).

#### 592 Phenotypic classification based on GSL content:

- 593 For each accession the expressed enzyme in each of the following families was determined based
- on the content (presence and amounts) of short chained Aliphatic GSLs:
- 595 MAM enzymes: the total amount of 3 carbons GSLs and 4 carbons GSLs was calculated for each
- accession. 3 carbons GSLs include 3MT, 3MSO, 3OHP and Allyl GSL. 4 carbons GSLs include
- 4MT, 4MSO, 4OHB, 3-butenyl and 2-OH-3-butenyl GSL (for structures and details see supp.
- Table 1). Accessions that the majority of Aliphatic short chained GSL contained 3 carbons in
- their side chains classified as MAM2 expressed (supp. Figure 3). Accessions that the majority of
- Aliphatic short chained GSL contained 4 carbons in their side chains classified as MAM1
- 601 expressed (supp. Figure 3). The accessions were plotted on a map based on their original
- 602 collection sites (supp. Figure 3).
- AOP enzymes: the relative amount of alkenyl GSL, alkyl GSL and MSO GSL were calculated in respect to the total short chained Aliphatic GSL as follows:

605 Alkenyl GSL (AOP2 expressed) = 
$$\frac{\text{Allyl}+2-\text{OH}-3-\text{butenyl}+3-\text{butenyl}}{\text{Total short chained GSL}}$$

- 606 Alkyl GSL (AOP3 expressed) =  $\frac{30HP + 40HB}{Total short chained GSL}$
- 607 MSO GSL (AOP null) =  $\frac{3MSO + 4MSO}{Total short chained GSL}$
- 608 The expressed AOP enzyme was determined based on those ratios: accessions with majority
- alkenyl GSL were classified as AOP2 expressed. Accessions with majority of alkyl GSL were
- classified as AOP3 expressed. Accessions with majority of MSO GSL were classified as AOP

- null. The accessions were plotted on a map based on their original collection sites (supp. Figure4).
- 613 GS-OH enzyme: the ratio between 2-OH-3-butenyl GSL to 3-butenyl GSL was calculated only
- for MAM1 expressed accessions (accessions that the majority of GSLs contain 4 carbons in their
- side chain). Accessions with high amounts of 2-OH-3-butenyl GSL were classified as GS-OH
- 616 functional. Accessions with high amounts of 3-butenyl GSL were classified as GS-OH non-
- functional. The accessions were plotted on a map based on their original collection sites (supp.
- 618 Figure 5).
- Each accession was classified to one of seven Aliphatic short chained GSLs based on the
- 620 combination of the dominancy of the enzymes as follows: MAM2, AOP null: classified as
- 621 3MSO dominant. MAM1, AOP null: classified as 4MSO dominant. MAM2, AOP3: classified as
- 622 3OHP dominant. MAM1, AOP3: classified as 4OHB dominant. MAM2, AOP2: classified as
- Allyl dominant. MAM1, AOP2, GS-OH non-functional: classified as 3-Butenyl dominant.
- MAM1, AOP2, GS-OH functional: classified as 2-OH-3-Butenyl dominant. The accessions were
- 625 plotted on a map based on their original collection sites and colored based on their dominant
- 626 chemotype (Figure 3).

# 627 Environmental data:

- Environmental data was obtained from the 1001 genomes website (<u>https://1001genomes.org/</u>, for
- 629 geographical data) and from the Arabidopsis CLIMtools
- 630 (<u>http://www.personal.psu.edu/sma3/CLIMtools.html</u>, (Ferrero-Serrano & Assmann, 2019)) for
- 631 environmental data. We used the five variables that captured a majority of the variance in this
- dataset including maximal temperature of warmest month (WC2\_BIO5), minimal temperature of
- coldest month (WC2\_BIO6), precipitation of wettest month (WC2\_BIO13), precipitation of
- driest month (WC2\_BIO14), and distance to the coast (in Km).

# 635 Environmental MANOVA:

- Linear models to test the effect of geographical and environmental parameters (supp. Figure 1, 8)
- 637 were conducted using dplyr package ("CRAN Package dplyr," n.d.) and included the following
- 638 parameters:

Supp. Figure 1- linear models for collection sites: PC score ~ Latitude + Longitude + Latitude \*
Longitude.

641 Supp. Figure 7 - for all the data: C length (C3 or C4) ~ Genomic group + Geography (north

642 versus south) +Max temperature of warmest month+ Min temperature of coldest month+

643 Precipitation of wettest month+ Precipitation of driest month+ Distance to the coast + Geography

\*Genomic group + Geography \* Max temperature of warmest month + Geography \* Min

temperature of coldest month+ Geography \* Precipitation of driest month+ Geography \*

646 Precipitation of wettest month + Geography \*Distance to the coast.

647 For the north and the south: C length (C3 or C4) ~ Genomic group + Geography (north versus

south) +Max temperature of warmest month+ Min temperature of coldest month+ Precipitation

of wettest month+ Precipitation of driest month+ Distance to the coast.

650 Multivariate analysis of variance (MANOVA) models to check the effect of environmental

variables on the chemotype identity for each collection (Figure 4 and supp. Figure 7) included

652 the following parameters:

Max temperature of warmest month+ Min temperature of coldest month+ Precipitation of wettest
month+ Precipitation of driest month+ Distance to the coast~ Chemotype.

**Random Forest analyses:** Random forest analyses was conducted using the "randomForest"

and "ElemStatLearn" packages in Rstudio ("CRAN - Package ElemStatLearn," n.d., "CRAN: R

News," n.d.; Liaw & Wiener, 2002). In these analyzes we used the environmental parameters

and genomic group data to predict the chemotype identity, after excluding the low frequencies

chemotypes (40HB and 3-Butenyl from all of them, 3MSO from the south).

660 Genome wide association studies:

The phenotypes for GWAS were each accession value for PC1 and 2. GWAS was implemented

with the easyGWAS tool (Grimm et al., 2017) using the EMMAX algorithms (Kang et al.,

663 2010) and a minor allele frequency (MAF) cutoff of 5%. The results were visualized as

664 manhattan plots using the qqman package in R (Turner, 2014).

665 **Phylogeny:** 

- 666 Genomic sequences from the accessions for MAM3 AT5G23020, AOP2 Chr4, 1351568 until
- 667 1354216, AOP3 AT4G03050.2, and GS-OH AT2G25450 were obtained using the
- 668 Pseudogenomes tool (<u>https://tools.1001genomes.org/pseudogenomes/#select\_strains</u>).
- 669 Multiple sequence alignment was done with the msa package in R, using the ClustalW,
- 670 ClustalOmega, and Muscle algorithms (Bodenhofer, Bonatesta, Horejš-Kainrath, & Hochreiter,
- 671 2015). Phylogenetic trees were generated with the ggtree package in R (Yu, 2020). Each tree was
- 672 rooted by the genes matching *Arabidopsis layrata's* functional orthologue or closest homologue.

673

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# 692 **Bibliography**

- Abdi, H., & Williams, L. J. (2010). Principal component analysis. *Wiley Interdisciplinary Reviews: Computational Statistics*, 2(4), 433–459. https://doi.org/10.1002/wics.101
- Abrahams, R. S., Pires, J. C., & Schranz, M. E. (2020). Genomic origin and diversification of the
  glucosinolate MAM locus. *Frontiers in Plant Science*, *11*, 711.
  https://doi.org/10.3389/fpls.2020.00711
- Agrawal, A. A. (2000). Overcompensation of plants in response to herbivory and the by-product
  benefits of mutualism. *Trends in Plant Science*, 5(7), 309–313.
  https://doi.org/10.1016/S1360-1385(00)01679-4
- Atwell, S., Huang, Y. S., Vilhjálmsson, B. J., Willems, G., Horton, M., Li, Y., ... Nordborg, M.
  (2010). Genome-wide association study of 107 phenotypes in Arabidopsis thaliana inbred lines. *Nature*, 465(7298), 627–631. https://doi.org/10.1038/nature08800
- Bakker, E. G., Traw, M. B., Toomajian, C., Kreitman, M., & Bergelson, J. (2008). Low levels of
   polymorphism in genes that control the activation of defense response in Arabidopsis
   thaliana. *Genetics*, 178(4), 2031–2043. https://doi.org/10.1534/genetics.107.083279
- Bednarek, P., & Osbourn, A. (2009). Plant-microbe interactions: chemical diversity in plant
   defense. *Science*, *324*(5928), 746–748. https://doi.org/10.1126/science.1171661
- Beekwilder, J., van Leeuwen, W., van Dam, N. M., Bertossi, M., Grandi, V., Mizzi, L., ... Bovy,
   A. (2008). The impact of the absence of aliphatic glucosinolates on insect herbivory in
   Arabidopsis. *Plos One*, *3*(4), e2068. https://doi.org/10.1371/journal.pone.0002068
- Benderoth, M., Textor, S., Windsor, A. J., Mitchell-Olds, T., Gershenzon, J., & Kroymann, J.
   (2006). Positive selection driving diversification in plant secondary metabolism.
   *Proceedings of the National Academy of Sciences of the United States of America*,
- 715 *103*(24), 9118–9123. https://doi.org/10.1073/pnas.0601738103
- Bialy, Z., Oleszek, W., Lewis, J., & Fenwick, G. R. (1990). Allelopathic potential of
  glucosinolates (mustard oil glycosides) and their degradation products against wheat. *Plant and Soil*, 129(2), 277–281. https://doi.org/10.1007/BF00032423
- Bodenhofer, U., Bonatesta, E., Horejš-Kainrath, C., & Hochreiter, S. (2015). msa: an R package
  for multiple sequence alignment. *Bioinformatics*, *31*(24), 3997–3999.
  https://doi.org/10.1093/bioinformatics/btv494
- Brachi, B., Meyer, C. G., Villoutreix, R., Platt, A., Morton, T. C., Roux, F., & Bergelson, J.
  (2015). Coselected genes determine adaptive variation in herbivore resistance throughout the native range of Arabidopsis thaliana. *Proceedings of the National Academy of Sciences of the United States of America*, 112(13), 4032–4037.
- 726 https://doi.org/10.1073/pnas.1421416112

727 728 729	Brown, P. D., Tokuhisa, J. G., Reichelt, M., & Gershenzon, J. (2003). Variation of glucosinolate accumulation among different organs and developmental stages of Arabidopsis thaliana. <i>Phytochemistry</i> , 62(3), 471–481. https://doi.org/10.1016/S0031-9422(02)00549-6
730	Chan, E. K. F., Rowe, H. C., Corwin, J. A., Joseph, B., & Kliebenstein, D. J. (2011). Combining
731	genome-wide association mapping and transcriptional networks to identify novel genes
732	controlling glucosinolates in Arabidopsis thaliana. <i>PLoS Biology</i> , 9(8), e1001125.
733	https://doi.org/10.1371/journal.pbio.1001125
734	Chan, E. K. F., Rowe, H. C., & Kliebenstein, D. J. (2010). Understanding the evolution of
735	defense metabolites in Arabidopsis thaliana using genome-wide association mapping.
736	<i>Genetics</i> , 185(3), 991–1007. https://doi.org/10.1534/genetics.109.108522
737	Corrion, A., & Day, B. (2001). Pathogen Resistance Signalling in Plants. In John Wiley & Sons
738	Ltd (Ed.), <i>eLS</i> (pp. 1–14). Chichester, UK: John Wiley & Sons, Ltd.
739	https://doi.org/10.1002/9780470015902.a0020119.pub2
740	CRAN - Package dplyr. (n.d.). Retrieved June 16, 2020, from https://CRAN.R-
741	project.org/package=dplyr
742	CRAN - Package ElemStatLearn. (n.d.). Retrieved June 16, 2020, from https://CRAN.R-
743	project.org/package=ElemStatLearn
744	CRAN - Package emmeans. (n.d.). Retrieved June 16, 2020, from https://CRAN.R-
745	project.org/package=emmeans
746	CRAN: R News. (n.d.). Retrieved June 16, 2020, from https://CRAN.R-project.org/doc/Rnews/
747 748 749	<ul> <li>Daxenbichler, M. E., Spencer, G. F., Carlson, D. G., Rose, G. B., Brinker, A. M., &amp; Powell, R. G. (1991). Glucosinolate composition of seeds from 297 species of wild plants. <i>Phytochemistry</i>, 30(8), 2623–2638. https://doi.org/10.1016/0031-9422(91)85112-D</li> </ul>
750	Erb, M., & Kliebenstein, D. J. (2020). Plant secondary metabolites as defenses, regulators, and
751	primary metabolites: the blurred functional trichotomy. <i>Plant Physiology</i> .
752	https://doi.org/10.1104/pp.20.00433
753	Fan, P., Leong, B. J., & Last, R. L. (2019). Tip of the trichome: evolution of acylsugar metabolic
754	diversity in Solanaceae. <i>Current Opinion in Plant Biology</i> , 49, 8–16.
755	https://doi.org/10.1016/j.pbi.2019.03.005
756	Ferrero-Serrano, Á., & Assmann, S. M. (2019). Phenotypic and genome-wide association with
757	the local environment of Arabidopsis. <i>Nature Ecology &amp; Evolution</i> , 3(2), 274–285.
758	https://doi.org/10.1038/s41559-018-0754-5
759	Futuyma, D. J., & Agrawal, A. A. (2009). Macroevolution and the biological diversity of plants
760	and herbivores. <i>Proceedings of the National Academy of Sciences of the United States of</i>
761	<i>America</i> , 106(43), 18054–18061. https://doi.org/10.1073/pnas.0904106106

762 763 764	<ul> <li>Giamoustaris, A., &amp; Mithen, R. (1996). Genetics of aliphatic glucosinolates. IV. Side-chain modification in Brassica oleracea. <i>TAG. Theoretical and Applied Genetics. Theoretische Und Angewandte Genetik</i>, 93(5-6), 1006–1010. https://doi.org/10.1007/BF00224105</li> </ul>
765	Grimm, D. G., Roqueiro, D., Salomé, P. A., Kleeberger, S., Greshake, B., Zhu, W.,
766	Borgwardt, K. M. (2017). easyGWAS: A Cloud-Based Platform for Comparing the
767	Results of Genome-Wide Association Studies. <i>The Plant Cell</i> , 29(1), 5–19.
768	https://doi.org/10.1105/tpc.16.00551
769	Halkier, B. A., & Gershenzon, J. (2006). Biology and biochemistry of glucosinolates. Annual
770	Review of Plant Biology, 57, 303–333.
771	https://doi.org/10.1146/annurev.arplant.57.032905.105228
772	Hanower, P., & Brzozowska, J. (1975). Influence d'un choc osmotique sur la composition des
773	feuilles de cotonnier en acides amines libres. <i>Phytochemistry</i> , 14(8), 1691–1694.
774	https://doi.org/10.1016/0031-9422(75)85275-7
775 776 777 778 779	<ul> <li>Hansen, B. G., Kerwin, R. E., Ober, J. A., Lambrix, V. M., Mitchell-Olds, T., Gershenzon, J.,</li> <li>Kliebenstein, D. J. (2008). A novel 2-oxoacid-dependent dioxygenase involved in the formation of the goiterogenic 2-hydroxybut-3-enyl glucosinolate and generalist insect resistance in Arabidopsis, <i>Plant Physiology</i>, <i>148</i>(4), 2096–2108. https://doi.org/10.1104/pp.108.129981</li> </ul>
780	Hansen, B. G., Kliebenstein, D. J., & Halkier, B. A. (2007). Identification of a flavin-
781	monooxygenase as the S-oxygenating enzyme in aliphatic glucosinolate biosynthesis in
782	Arabidopsis. <i>The Plant Journal: For Cell and Molecular Biology</i> , 50(5), 902–910.
783	https://doi.org/10.1111/j.1365-313X.2007.03101.x
784	Hasegawa, T., Yamada, K., Kosemura, S., Yamamura, S., & Hasegawa, K. (2000). Phototropic
785	stimulation induces the conversion of glucosinolate to phototropism-regulating
786	substances of radish hypocotyls. <i>Phytochemistry</i> , 54(3), 275–279.
787	https://doi.org/10.1016/S0031-9422(00)00080-7
788	Hayat, S., Hayat, Q., Alyemeni, M. N., Wani, A. S., Pichtel, J., & Ahmad, A. (2012). Role of
789	proline under changing environments: a review. <i>Plant Signaling &amp; Behavior</i> , 7(11),
790	1456–1466. https://doi.org/10.4161/psb.21949
791	Heidel, A. J., Clauss, M. J., Kroymann, J., Savolainen, O., & Mitchell-Olds, T. (2006). Natural
792	variation in MAM within and between populations of Arabidopsis lyrata determines
793	glucosinolate phenotype. <i>Genetics</i> , 173(3), 1629–1636.
794	https://doi.org/10.1534/genetics.106.056986
795 796	https://journal.r-project.org/archive/2013-1/kahle-wickham.pdf. (n.d.). Retrieved June 16, 2020, from https://journal.r-project.org/archive/2013-1/kahle-wickham.pdf
797	Hu, L., Mateo, P., Ye, M., Zhang, X., Berset, J. D., Handrick, V., Erb, M. (2018). Plant iron
798	acquisition strategy exploited by an insect herbivore. <i>Science</i> , 361(6403), 694–697.
799	https://doi.org/10.1126/science.aat4082

800 801 802	Jander, G., Cui, J., Nhan, B., Pierce, N. E., & Ausubel, F. M. (2001). The TASTY locus on chromosome 1 of Arabidopsis affects feeding of the insect herbivore Trichoplusia ni. <i>Plant Physiology</i> , 126(2), 890–898. https://doi.org/10.1104/pp.126.2.890
803 804	Kahle, D., & Wickham, H. (2013). ggmap: Spatial Visualization with ggplot2. <i>The R Journal</i> , 5(1), 144. https://doi.org/10.32614/RJ-2013-014
805 806 807	<ul> <li>Kang, H. M., Sul, J. H., Service, S. K., Zaitlen, N. A., Kong, SY., Freimer, N. B., Eskin, E. (2010). Variance component model to account for sample structure in genome-wide association studies. <i>Nature Genetics</i>, 42(4), 348–354. https://doi.org/10.1038/ng.548</li> </ul>
808 809 810 811	Katz, E., Bagchi, R., Jeschke, V., Rasmussen, A. R. M., Hopper, A., Burow, M., Kliebenstein, D. J. (2020). Diverse allyl glucosinolate catabolites independently influence root growth and development. <i>Plant Physiology</i> , 183(3), 1376–1390. https://doi.org/10.1104/pp.20.00170
812 813 814	Katz, E., Nisani, S., Sela, M., Behar, H., & Chamovitz, D. A. (2015). The effect of indole-3- carbinol on PIN1 and PIN2 in Arabidopsis roots. <i>Plant Signaling &amp; Behavior</i> , 10(9), e1062200. https://doi.org/10.1080/15592324.2015.1062200
815 816 817 818	<ul> <li>Kerwin, R. E., Feusier, J., Muok, A., Lin, C., Larson, B., Copeland, D., Kliebenstein, D. J. (2017). Epistasis × environment interactions among Arabidopsis thaliana glucosinolate genes impact complex traits and fitness in the field. <i>The New Phytologist</i>, 215(3), 1249–1263. https://doi.org/10.1111/nph.14646</li> </ul>
819 820 821	Kerwin, R., Feusier, J., Corwin, J., Rubin, M., Lin, C., Muok, A., Kliebenstein, D. J. (2015). Natural genetic variation in Arabidopsis thaliana defense metabolism genes modulates field fitness. <i>eLife</i> , <i>4</i> . https://doi.org/10.7554/eLife.05604
822 823 824 825	Kim, J., Kang, K., Gonzales-Vigil, E., Shi, F., Jones, A. D., Barry, C. S., & Last, R. L. (2012). Striking natural diversity in glandular trichome acylsugar composition is shaped by variation at the Acyltransferase2 locus in the wild tomato Solanum habrochaites. <i>Plant</i> <i>Physiology</i> , <i>160</i> (4), 1854–1870. https://doi.org/10.1104/pp.112.204735
826 827 828	Kliebenstein, D J. (2004). Secondary metabolites and plant/environment interactions: a view through Arabidopsis thaliana tinged glasses. <i>Plant, Cell &amp; Environment</i> , 27(6), 675–684. https://doi.org/10.1111/j.1365-3040.2004.01180.x
829 830 831	<ul> <li>Kliebenstein, D J, &amp; Cacho, N. I. (2016). Nonlinear Selection and a Blend of Convergent, Divergent and Parallel Evolution Shapes Natural Variation in Glucosinolates. In <i>In S.</i> <i>Kopriva (Ed.), Glucosinolates</i>. Elsevier. https://doi.org/10.1016/bs.abr.2016.06.002</li> </ul>
832 833 834	Kliebenstein, D J, Gershenzon, J., & Mitchell-Olds, T. (2001). Comparative quantitative trait loci mapping of aliphatic, indolic and benzylic glucosinolate production in Arabidopsis thaliana leaves and seeds. <i>Genetics</i> , <i>159</i> (1), 359–370.
835 836	Kliebenstein, D J, Kroymann, J., Brown, P., Figuth, A., Pedersen, D., Gershenzon, J., & Mitchell-Olds, T. (2001). Genetic control of natural variation in Arabidopsis

837	glucosinolate accumulation. <i>Plant Physiology</i> , 126(2), 811–825.
838	https://doi.org/10.1104/pp.126.2.811
839 840 841 842	<ul> <li>Kliebenstein, D J, Lambrix, V. M., Reichelt, M., Gershenzon, J., &amp; Mitchell-Olds, T. (2001).</li> <li>Gene duplication in the diversification of secondary metabolism: tandem 2-oxoglutarate- dependent dioxygenases control glucosinolate biosynthesis in Arabidopsis. <i>The Plant</i> <i>Cell</i>, 13(3), 681–693. https://doi.org/10.1105/tpc.13.3.681</li> </ul>
843 844 845	Kliebenstein, D., Pedersen, D., Barker, B., & Mitchell-Olds, T. (2002). Comparative analysis of quantitative trait loci controlling glucosinolates, myrosinase and insect resistance in Arabidopsis thaliana. <i>Genetics</i> , <i>161</i> (1), 325–332.
846 847	Kliebenstein, Daniel J, Figuth, A., & Mitchell-Olds, T. (2002). Genetic architecture of plastic methyl jasmonate responses in Arabidopsis thaliana. <i>Genetics</i> , <i>161</i> (4), 1685–1696.
848	Kliebenstein, Daniel J. (2009). A quantitative genetics and ecological model system:
849	understanding the aliphatic glucosinolate biosynthetic network via QTLs. <i>Phytochemistry</i>
850	<i>Reviews : Proceedings of the Phytochemical Society of Europe</i> , 8(1), 243–254.
851	https://doi.org/10.1007/s11101-008-9102-8
852	Kroymann, J., Donnerhacke, S., Schnabelrauch, D., & Mitchell-Olds, T. (2003). Evolutionary
853	dynamics of an Arabidopsis insect resistance quantitative trait locus. <i>Proceedings of the</i>
854	<i>National Academy of Sciences of the United States of America</i> , 100 Suppl 2, 14587–
855	14592. https://doi.org/10.1073/pnas.1734046100
856	Kroymann, J., & Mitchell-Olds, T. (2005). Epistasis and balanced polymorphism influencing
857	complex trait variation. <i>Nature</i> , 435(7038), 95–98. https://doi.org/10.1038/nature03480
858 859 860	Lankau, R. A. (2007). Specialist and generalist herbivores exert opposing selection on a chemical defense. <i>The New Phytologist</i> , <i>175</i> (1), 176–184. https://doi.org/10.1111/j.1469-8137.2007.02090.x
861	Lankau, R. A., & Kliebenstein, D. J. (2009). Competition, herbivory and genetics interact to
862	determine the accumulation and fitness consequences of a defence metabolite. <i>Journal of</i>
863	<i>Ecology</i> , 97(1), 78–88. https://doi.org/10.1111/j.1365-2745.2008.01448.x
864	Lankau, R. A., & Strauss, S. Y. (2007). Mutual feedbacks maintain both genetic and species
865	diversity in a plant community. <i>Science</i> , 317(5844), 1561–1563.
866	https://doi.org/10.1126/science.1147455
867	Lankau, R. A., & Strauss, S. Y. (2008). Community complexity drives patterns of natural
868	selection on a chemical defense of Brassica nigra. <i>The American Naturalist</i> , 171(2), 150–
869	161. https://doi.org/10.1086/524959
870	Liaw, A., & Wiener, M. (2002). Classification and Regression by randomForest. R News 2.
871 872 873	MacQueen, A., Sun, X., & Bergelson, J. (2016). Genetic architecture and pleiotropy shape costs of Rps2-mediated resistance in Arabidopsis thaliana. <i>Nature Plants</i> , <i>2</i> , 16110. https://doi.org/10.1038/nplants.2016.110

874 875	Malcolm, S. B. (1994). Milkweeds, monarch butterflies and the ecological significance of cardenolides. <i>Chemoecology</i> , <i>5-6</i> (3-4), 101–117. https://doi.org/10.1007/BF01240595
876 877 878 879	<ul> <li>Malinovsky, F. G., Thomsen, ML. F., Nintemann, S. J., Jagd, L. M., Bourgine, B., Burow, M., &amp; Kliebenstein, D. J. (2017). An evolutionarily young defense metabolite influences the root growth of plants via the ancient TOR signaling pathway. <i>eLife</i>, <i>6</i>. https://doi.org/10.7554/eLife.29353</li> </ul>
880	Mithen, R., Clarke, J., Lister, C., & Dean, C. (1995). Genetics of aliphatic glucosinolates. III.
881	Side chain structure of aliphatic glucosinolates in Arabidopsis thaliana. <i>Heredity</i> , 74(2),
882	210–215. https://doi.org/10.1038/hdy.1995.29
883 884 885 886	<ul> <li>Moore, B. M., Wang, P., Fan, P., Leong, B., Schenck, C. A., Lloyd, J. P., Shiu, SH. (2019).</li> <li>Robust predictions of specialized metabolism genes through machine learning.</li> <li><i>Proceedings of the National Academy of Sciences of the United States of America</i>, 116(6), 2344–2353. https://doi.org/10.1073/pnas.1817074116</li> </ul>
887 888	Opitz, S. E. W., & Müller, C. (2009). Plant chemistry and insect sequestration. <i>Chemoecology</i> , <i>19</i> (3), 117–154. https://doi.org/10.1007/s00049-009-0018-6
889	Petersen, B. L., Chen, S., Hansen, C. H., Olsen, C. E., & Halkier, B. A. (2002). Composition and
890	content of glucosinolates in developing Arabidopsis thaliana. <i>Planta</i> , 214(4), 562–571.
891	https://doi.org/10.1007/s004250100659
892 893 894 895	Pfalz, M., Vogel, H., Mitchell-Olds, T., & Kroymann, J. (2007). Mapping of QTL for resistance against the crucifer specialist herbivore <i>Pieris brassicae</i> in a new Arabidopsis inbred line population, Da(1)-12 x Ei-2. <i>Plos One</i> , 2(6), e578. https://doi.org/10.1371/journal.pone.0000578
896	Ramos-Onsins, S. E., Stranger, B. E., Mitchell-Olds, T., & Aguadé, M. (2004). Multilocus
897	Analysis of Variation and Speciation in the Closely Related Species Arabidopsis halleri
898	and A. lyrata. Genetics, 166(1), 373–388. https://doi.org/10.1534/genetics.166.1.373
899	Raybould, A. F., & Moyes, C. L. (2001). The ecological genetics of aliphatic glucosinolates.
900	<i>Heredity</i> , 87(Pt 4), 383–391. https://doi.org/10.1046/j.1365-2540.2001.00954.x
901	Rodman, James E., Kruckeberg, A. R., & Al-Shehbaz, I. A. (1981). Chemotaxonomic diversity
902	and complexity in seed glucosinolates of caulanthus and streptanthus (cruciferae).
903	<i>Systematic Botany</i> , 6(3), 197. https://doi.org/10.2307/2418282
904 905 906	Rodman, James Eric. (1980). Population variation and hybridization in sea-rockets (cakile, cruciferae): seed glucosinolate characters. <i>American Journal of Botany</i> , 67(8), 1145–1159. https://doi.org/10.1002/j.1537-2197.1980.tb07748.x
907	Salehin, M., Li, B., Tang, M., Katz, E., Song, L., Ecker, J. R., Estelle, M. (2019). Auxin-
908	sensitive Aux/IAA proteins mediate drought tolerance in Arabidopsis by regulating
909	glucosinolate levels. <i>Nature Communications</i> , 10(1), 4021.
910	https://doi.org/10.1038/s41467-019-12002-1

911	Schilmiller, A. L., Pichersky, E., & Last, R. L. (2012). Taming the hydra of specialized
912	metabolism: how systems biology and comparative approaches are revolutionizing plant
913	biochemistry. <i>Current Opinion in Plant Biology</i> , 15(3), 338–344.
914	https://doi.org/10.1016/j.pbi.2011.12.005
915	Sønderby, I. E., Geu-Flores, F., & Halkier, B. A. (2010). Biosynthesis of glucosinolatesgene
916	discovery and beyond. <i>Trends in Plant Science</i> , 15(5), 283–290.
917	https://doi.org/10.1016/j.tplants.2010.02.005
918	Szakiel, A., Pączkowski, C., & Henry, M. (2011). Influence of environmental biotic factors on
919	the content of saponins in plants. <i>Phytochemistry Reviews : Proceedings of the</i>
920	<i>Phytochemical Society of Europe</i> , 10(4), 493–502. https://doi.org/10.1007/s11101-010-
921	9164-2
922 923 924	Thakur, P., & Rai, V. (1982). Dynamics of amino acid accumulation of two differentially drought resistant Zea mays cultivars in response to osmotic stress. <i>Environmental and Experimental Botany</i> , 22(2), 221–226. https://doi.org/10.1016/0098-8472(82)90042-9
925 926 927	The 1001 Genomes Consortium. (2016). 1,135 Genomes reveal the global pattern of polymorphism in <i>Arabidopsis thaliana</i> . <i>Cell</i> , <i>166</i> (2), 481–491. https://doi.org/10.1016/j.cell.2016.05.063
928	Todesco, M., Balasubramanian, S., Hu, T. T., Traw, M. B., Horton, M., Epple, P., Weigel, D.
929	(2010). Natural allelic variation underlying a major fitness trade-off in Arabidopsis
930	thaliana. <i>Nature</i> , 465(7298), 632–636. https://doi.org/10.1038/nature09083
931 932	Turner, S. D. (2014). qqman: an R package for visualizing GWAS results using Q-Q and manhattan plots. <i>BioRxiv</i> . https://doi.org/10.1101/005165
933	Uremis, I., Arslan, M., Sangun, M. K., Uygur, V., & Isler, N. (2009). Allelopathic potential of
934	rapeseed cultivars on germination and seedling growth of weeds. <i>Asian Journal of</i>
935	<i>Chemistry</i> , 21(3), 2170–2184.
936	Wentzell, A. M., & Kliebenstein, D. J. (2008). Genotype, age, tissue, and environment regulate
937	the structural outcome of glucosinolate activation. <i>Plant Physiology</i> , 147(1), 415–428.
938	https://doi.org/10.1104/pp.107.115279
939	Wentzell, A. M., Rowe, H. C., Hansen, B. G., Ticconi, C., Halkier, B. A., & Kliebenstein, D. J.
940	(2007). Linking metabolic QTLs with network and cis-eQTLs controlling biosynthetic
941	pathways. <i>PLoS Genetics</i> , 3(9), 1687–1701.
942	https://doi.org/10.1371/journal.pgen.0030162
943	Wickham, H., Averick, M., Bryan, J., Chang, W., McGowan, L., François, R., Yutani, H.
944	(2019). Welcome to the tidyverse. <i>The Journal of Open Source Software</i> , 4(43), 1686.
945	https://doi.org/10.21105/joss.01686
946	Windsor, A. J., Reichelt, M., Figuth, A., Svatos, A., Kroymann, J., Kliebenstein, D. J.,
947	Mitchell-Olds, T. (2005). Geographic and evolutionary diversification of glucosinolates

948 949	among near relatives of Arabidopsis thaliana (Brassicaceae). <i>Phytochemistry</i> , 66(11), 1321–1333. https://doi.org/10.1016/j.phytochem.2005.04.016
950	Wolters, H., & Jürgens, G. (2009). Survival of the flexible: hormonal growth control and
951	adaptation in plant development. <i>Nature Reviews. Genetics</i> , 10(5), 305–317.
952	https://doi.org/10.1038/nrg2558
953	Wright, S. I., Lauga, B., & Charlesworth, D. (2002). Rates and patterns of molecular evolution in
954	inbred and outbred Arabidopsis. <i>Molecular Biology and Evolution</i> , 19(9), 1407–1420.
955	https://doi.org/10.1093/oxfordjournals.molbev.a004204
956	Yamada, K., Hasegawa, T., Minami, E., Shibuya, N., Kosemura, S., Yamamura, S., &
957	Hasegawa, K. (2003). Induction of myrosinase gene expression and myrosinase activity
958	in radish hypocotyls by phototropic stimulation. <i>Journal of Plant Physiology</i> , 160(3),
959	255–259. https://doi.org/10.1078/0176-1617-00950
960	Yang, C. W., Lin, C. C., & Kao, C. H. (2000). Proline, ornithine, arginine and glutamic acid
961	contents in detached rice leaves. <i>Biologia Plantarum</i> , 43(2), 305–307.
962	https://doi.org/10.1023/A:1002733117506
963 964	Yu, G. (2020). Using ggtree to Visualize Data on Tree-Like Structures. <i>Current Protocols in Bioinformatics</i> , 69(1), e96. https://doi.org/10.1002/cpbi.96
965	Zhu, W., Zaidem, M., Van deaaa 1Weyer, AL., Gutaker, R. M., Chae, E., Kim, ST.,
966	Weigel, D. (2018). Modulation of ACD6 dependent hyperimmunity by natural alleles of
967	an Arabidopsis thaliana NLR resistance gene. <i>PLoS Genetics</i> , 14(9), e1007628.
968	https://doi.org/10.1371/journal.pgen.1007628
969	Züst, T., & Agrawal, A. A. (2017). Trade-Offs Between Plant Growth and Defense Against
970	Insect Herbivory: An Emerging Mechanistic Synthesis. <i>Annual Review of Plant Biology</i> ,
971	68, 513–534. https://doi.org/10.1146/annurev-arplant-042916-040856
972	Züst, T., Heichinger, C., Grossniklaus, U., Harrington, R., Kliebenstein, D. J., & Turnbull, L. A.
973	(2012). Natural enemies drive geographic variation in plant defenses. <i>Science</i> , 338(6103),
974	116–119. https://doi.org/10.1126/science.1226397
975	