An integrative approach to investigate natural variation

² in the accumulation of aliphatic glucosinolates in

3 Arabidopsis thaliana

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10 Abstract

11 Glucosinolates are a fascinating class of specialised metabolites found in the plants of Brassicacea family. The variation in glucosinolate composition across different Arabidopsis 12 13 ecotypes could be a result of allelic compositions at different biosynthetic loci. The 14 contribution of methylthioalkylmalate synthase (MAM) genes to diversity of glucosinolate profiles across different Arabidopsis ecotypes has been confirmed by genetic analyses. 15 16 Different MAM isoforms utilise different chain-elongated substrates for glucosinolate 17 biosynthesis causing thus a variation in chain lengths across different Arabidopsis ecotypes. 18 To further investigate the relationship between the genotype and the associated metabolic 19 phenotype, we studied the diversity of genes and enzymes of glucosinolate biosynthesis. 20 Using Shannon entropy as a measure we revealed that several genes of the pathway show a 21 clear derivation from the expected behaviour, either accumulating non-synonymous SNPs or 22 showing signs of purifying selection. We found that the genotype-phenotype relationship is 23 much more complicated than inferred from the diversity of MAM synthases. We conclude 24 therefore, that the ON/OFF feature of key QTLs is not enough to elucidate the diversity of 25 glucosinolates across different Arabidopsis thaliana ecotypes and that glucosinolate profiles 26 are determined also through the polymorphic residues along the coding regions of multiple metabolic genes. 27

28 Introduction

29 Plant secondary metabolism produces a huge variation in structures and molecules with a 30 plethora of functions for the plants but also for human nutrition and health (Owen, Patron, 31 Huang, & Osbourn, 2017). One class of such secondary metabolites are glucosinolates in the 32 Brassicaceae. Glucosinolates (GSLs) are important for the plants as a defence against 33 herbivores, fungi, and other pathogens (Halkier & Gershenzon, 2006). They are also 34 determinants of taste and flavour of cruciferous vegetables and responsible for their positive 35 health properties (Traka & Mithen, 2009). More than 140 different GSL structures have been 36 described, with a great variation not only between species, but also among ecotypes of the 37 same species (Agerbirk et al., 2015; Clarke, 2010). GSLs are synthesised from amino acids 38 and are divided into three classes: aliphatic GSLs, derived from alanine, leucine, isoleucine, 39 valine and methionine (Met), indolic GSLs synthesised from tryptophan, and aromatic GSLs 40 from phenylalanine (Phe) (Halkier & Gershenzon, 2006; Sønderby, Geu-Flores, & Halkier, 41 2010). All GSLs possess the same core structure, which comprises a glucose residue linked to 42 a (Z)-N-hydroximic sulfate ester via a sulphur atom (Fahey, Zalcmann, & Talalay, 2001). 43 GSL biosynthesis consists of three independent steps: (i) chain elongation of selected 44 precursor amino acids (only Met and Phe), (ii) formation of the core GSL structure, and (iii) 45 secondary modifications of the amino acid side chain. The diversity of GSLs derives from the 46 side-chain elongation and secondary modifications.

47 GSLs are best known for their function in interaction between plants and herbivores. 48 Upon tissue damage, the GSLs stored in the vacuoles come in contact with the enzyme 49 myrosinase, which cleaves the thio-glucoside bond. The resulting aglycones are unstable and 50 form volatile isothiocyanates or nitriles (Halkier & Gershenzon, 2006). Depending upon the 51 herbivore, the volatiles of specific GSLs can act as feeding deterrents or stimulants (Buskov, 52 Serra, Rosa, Sørensen, & Sørensen, 2002; Gabrys & Tjallingii, 2002; Lazzeri, Curto, Leoni, 53 & Dallavalle, 2004; Mewis, Ulrich, & Schnitzler, 2002; Miles, Campo, & Renwick, 2005; Noret et al., 2005). Therefore, there is often an overlap between quantitative trait loci (QTL) 54 55 for GSL accumulation and insect resistance (Kroymann, Donnerhacke, Schnabelrauch, & 56 Mitchell-Olds, 2003). A possible outcome of this heterogeneous natural selection on GSLs is 57 the quick evolution of new compounds or new patterns of compound accumulation 58 (Daxenbichler et al., 1991; Rodman, 1980). New GSLs may increase resistance to herbivores 59 that have become adapted to existing defences, whereas new patterns of GSL accumulation

may provide a unique complement of defences by slowing down the counter-adaptation ofherbivores.

62 The GSL defence system is one of the few systems where systematic data assessing 63 between and within species variation at both phenotypic and causal genetic level is available 64 (Halkier & Gershenzon, 2006; Sønderby et al., 2010). Natural variation within or between 65 species is regulated by a complex network of genes and associated polymorphisms (Fisher, 66 1930; Kliebenstein, Kroymann, et al., 2001; Lynch, Walsh, & others, 1998). These variations, 67 however, complicate our understanding of how certain genes behave in context of a species 68 as we often study a single genotype. Thus, understanding a metabolic pathway requires 69 studies involving more than one ecotype. For example, methionine derived aliphatic GSLs 70 differ in length of the side chain caused by the elongation of the amino acid, as well as by 71 further modifications, e.g. oxidation of the sulfur atom (Halkier & Gershenzon, 2006; 72 Sønderby et al., 2010). In the model plant Arabidopsis thaliana, aliphatic GSLs of six 73 different chain-lengths, referred to as 3C to 8C GSLs, but with different side chains are 74 found. The diversity in length of aliphatic GSLs can be explained by the variation in the 75 iterative chain-elongation cycles, each adding one methylene group to the Met and/or 76 elongated Met molecule (Halkier & Gershenzon, 2006). The QTL responsible for 77 determining the chain-elongation of GSLs is GS-ELONG (Magrath et al., 1994). GS-ELONG 78 is highly variable across different Arabidopsis ecotypes, with common large indel 79 polymorphism (Kroymann et al., 2003). The gene underlying the GS-ELONG QTL is 80 methylthioalkylmalate synthase (MAM3), encoding the key enzyme of the chain elongation 81 cycle (Kroymann et al., 2001). However, the GS-ELONG locus harbours two more genes, 82 isoforms of MAM3 called MAM1 and MAM2. MAM3 is present in all Arabidopsis ecotypes, 83 and some ecotypes possess both additional genes whereas others possess either MAM1 or 84 MAM2, with some, such as Landsberg erecta (Ler), having a truncated (non-functional) 85 MAM1 in addition to MAM2 (Kroymann et al., 2003). While the presence of functional 86 MAM genes has been described as responsible for the variation in aliphatic GSL side chain 87 length, very little is known about contribution of other genetic variation, mainly single 88 nucleotide polymorphism (SNP), at this locus.

In this paper, we investigate the link between the diversity of GSL enzyme-coding genes and their GSL profiles exhibited across 72 different *A. thaliana* ecotypes. The selection of 72 ecotypes is based on the availability of information about the gene sequences and the patterns of accumulation of aliphatic glucosinolates. Importantly, the experiments for

93 determining the aliphatic glucosinolate levels have been performed under identical conditions 94 (Chan, Rowe, & Kliebenstein, 2010; Kliebenstein, Kroymann, et al., 2001). It can therefore 95 be assumed that the environment was identical (up to experimental precision) for all 96 ecotypes. This study presents an effort to quantify the impact of the diversity of MAM gene 97 sequence on GSL variation, rather than the on/off nature of the GS ELONG QTLs. The 98 genetic sequence coding for an enzyme determines the kinetic properties of the corresponding 99 enzyme. For example, polymorphisms in the active sites, in principle, can change the 100 substrate specificity of the respective enzyme. Thus, we investigate the level of 101 polymorphisms in the coding region of the GSL enzymes to study the impact on the diversity 102 of aliphatic GSLs.

103 Results

104 Distribution of MAM genes across Arabidopsis thaliana ecotypes

105 The genetic basis of chain-length distribution of aliphatic GSLs became evident with the 106 identification of the GS-ELONG QTL in Arabidopsis and Brassica napus (Magrath et al., 107 1994). The locus was mapped in Arabidopsis by using a cross between two ecotypes, 108 Columbia (Col) and Landsberg erecta (Ler), where the major glucosinolates are 4C and 3C, 109 respectively (Kroymann et al., 2001). The underlying candidates MAM1 and MAM3 genes are 110 two adjacent sequences with high similarity to genes encoding isopropylmalate synthase that 111 catalyses the condensation of chain elongation in leucine biosynthesis. Later, a third MAM-112 like gene, MAM2, was identified at the same locus as MAM1 (Kroymann et al., 2003). While 113 MAM3 is ubiquitous, most Arabidopsis ecotypes examined possessed functional copies of 114 either MAM1 or MAM2 genes. A functional MAM1 gene has been correlated with the 115 accumulation of 4C GSLs, whereas a functional MAM2 has been linked to 3C GSLs. To gain 116 more insights on the impact of MAM synthases on chain lengths distribution of aliphatic 117 GSL, we analysed the similarity of the annotated *MAM1* gene across 72 Arabidopsis ecotypes 118 taken from 1001 genome project database (Jorge et al., 2016). These ecotypes were selected 119 based on the availability of gene sequences and the associated aliphatic GSL profiles. Details 120 on the 72 ecotypes are given in the Supplementary Table 1. The annotated MAM1 sequences 121 from the 72 ecotypes were compared to each other for diversity. Figure 1 shows a mid-point 122 rooted phylogenetic tree showing the evolutionary relationship between the ecotypes based 123 on the similarities and differences in the coding region of the MAM1/MAM2 sequences. 124 Based on maximum likelihood estimation (Guindon et al., 2010), the tree shows two main

branches. While 53 out of the 72 ecotypes clustering in the blue branch indeed possess high

similarity to the coding region of MAM1 gene, 19 ecotypes in the red branch possess genes

127 more like the MAM2 gene. Thus, we assume that the ecotypes composed in blue and red

128 branches possess *MAM1* and *MAM2* genes, respectively.

129 The metabolic genotypes and associated phenotypes

130 We define a metabolic genotype (G_i) as the gene sequence of enzymes of glucosinolate 131 biosynthesis in ecotype i, whereas the metabolic phenotype (P_i) corresponds to the 132 composition of aliphatic GSLs in the ecotype *i*. To gain a deeper understanding of how 133 different metabolic genotypes and their associated phenotypes are linked, we analysed the genotypic and phenotypic distances. The genotypic distances between the genotypes were 134 calculated as Hamming distance (Hamming, 1950) $d_{i,i}^{G}$, which is the number of positions at 135 which the corresponding nucleotide/amino-acid characters are different between gene 136 sequences G_i and G_j of equal length. The phenotypic distance $d_{i,j}^{P}$ was calculated as 137 Euclidean distance, $d_{i,j}^P = \sqrt{\sum_{k=1}^n (P_{i,k} - P_{j,k})^2}$ between two phenotypes $P_{i,k} =$ 138 $(P_{i,1}, P_{i,2}, \dots, P_{i,n})$ and $P_{j,k} = (P_{j,1}, P_{j,2}, \dots, P_{j,n})$ in an *n*-dimensional space. In this study, 139 140 n=6, which corresponds to the total number of chain-elongated aliphatic GSLs found in A. thaliana (Figure 2). Thus, we can quantify differences between all pairs of ecotypes based on 141 their metabolic genotype G_i (i = 1, ..., 72) and phenotype P_i (i = 1, ..., 72). Figure 3 142 143 showcases the summary of the analysis, where the genotypic distances are plotted against the 144 phenotypic distances. Intuitively, one would assume that similar genotypes shall exhibit 145 similar phenotypes, and *vice-versa*. However, Figure 3 clearly shows that several ecotypes 146 show low genotypic distance (i.e. they are genotypically similar) but exhibit high phenotypic 147 distance (variation in GSL profiles). Also, there exist genotypically diverse ecotypes that 148 exhibit very similar GSL profiles. Thus, investigating the factors affecting variations in the 149 phenotypes of such ecotypes will provide a clearer understanding of how distinct patterns of GSL accumulation emerge out of genetic differences. Moreover, investigation of the 150 151 localisation of polymorphic residues in the GSL biosynthesis enzymes will provide a better understanding of the link between metabolic genotype and the associated metabolic 152 153 phenotypes.

154 Diversity of GSL enzyme-coding genes

155 To investigate the diversity of metabolic genes of GSL biosynthesis, we investigated the 156 levels of amino acid and nucleotide polymorphism across the 72 A. thaliana ecotypes by 157 calculating the average Shannon entropy (Shannon) H across the gene length (Figure 4A and 158 B). The analysis revealed that some of these enzymes are highly diverse while others remain 159 conserved across different ecotypes. Interestingly, the diversity seems to be independent of 160 steps of GSL biosynthesis in which the enzymes are active. From the diversity of amino acid 161 sequences (Figure 4A), FMO-GSOX1 enzyme exhibits the highest diversity (entropy), while 162 the lowest diversity is found in SOT17. Among the enzymes active in the chain-elongation 163 pathway, MAM1 shows the highest diversity while BAT5 shows the low diversity. However, a 164 further analysis of the diversity in the nucleotide sequences of the metabolic genotypes 165 showed a high level of polymorphism in BAT5 (cf. Figure 4B), which was not reflected in 166 the diversity of amino acid sequences. Indeed, most genes show only a slightly lower 167 diversity in amino acid variation than nucleotide variation, which reflects the degeneration of 168 the genetic code (Figure 4C). A plausible explanation for the low amino acid variation in 169 BAT5 could be the specificity towards a variety of chain-elongated substrates of GSL 170 biosynthesis (Halkier & Gershenzon, 2006). The low diversity of BAT5 could be linked to its 171 function in transport of a diverse range of compounds that are a part of aliphatic GSL 172 biosynthesis and Met-salvage pathway (Gigolashvili et al., 2009; Sauter, Moffatt, Saechao, 173 Hell, & Wirtz, 2013), thus, mutations in the coding region of BAT5 may impair the 174 functioning of both pathways. In contrast to BAT5, FMO-GSOX1 shows a low diversity in the 175 nucleotide sequences of 72 genotypes but reflect a high diversity in the amino acid 176 sequences. This is a clear example of preferential accumulation of non-synonymous 177 mutations, which alter the amino acid sequence of an enzyme.

High diversity of *MAM1* could be a consequence of incorrect annotation of *MAM1/MAM2*enzymes across 72 *A. thaliana* ecotypes. Thus, we analysed the diversity of GSL enzymes
across the MAM1 ecotypes (blue branch of Figure 1) and MAM2 (red branch of Figure 1)
ecotypes, separately. We did see a reduction in the diversity of MAM1 and MAM2 enzymes
(cf. Supplementary Figure 1 and Supplementary Figure 2). Nevertheless, *MAM1* and *MAM2*are still the most diverse enzymes of chain-elongation pathway.

184 Polymorphisms in the active sites of MAM enzymes

To get a clearer understanding of the effects of the localisation of polymorphic amino acid residues in the active sites of the metabolic enzymes, we extracted the information about the 187 active sites from the NCBI's conserved domain database (Marchler-Bauer et al., 2015). For 188 example, the amino acid positions 93, 94, 97, 124, 162, 164, 186, 227, 229, 231, 257, 259, 189 260, 261, 262, 290, 292, and 294 are known to be key for activity of MAM synthases, based 190 on the database and a crystal structure (Kumar et al., 2019; Marchler-Bauer et al., 2015; 191 Petersen et al., 2019). We refer to the amino acid positions from 93 to 294 as active region of 192 the enzyme. We have found that MAM synthases exhibit a maximum of 13 and 3 193 polymorphic residues in the active region of MAM1 and MAM3, respectively. Figure 5(A)194 and (B) show pairwise comparisons of polymorphisms in the active region of MAM 195 synthases against the genotypic distances of 72 A. thaliana ecotypes. Furthermore, we 196 recorded the polymorphisms at different positions in the active region of MAM synthases 197 (see Figure 5(C)). The amino acid residues at position 98, 99, 132, 138, 139, 147, 165, 173, 198 177, 187, 228, 245, 257, 258, 271, 289 and 290 of the MAM1 and positions 156, 231, 241 and 199 242 of MAM3 accumulate polymorphic residues across the 72 Arabidopsis ecotypes. 200 Polymorphisms in the active sites of an enzyme, in principle, can change the catalytic 201 properties of the enzyme (Kroymann et al., 2001; Kumar et al., 2019; Petersen et al., 2019). 202 However, the quantitative effect on the enzymatic properties cannot be explained due to 203 unavailability of enzyme abundances in these 72 ecotypes.

204 Discussion

205 Plasticity of the metabolic genotype and the associated GSL profiles

206 Glucosinolate metabolism results in a highly variable composition of individual metabolites 207 in Arabidopsis accessions, which is reflected by a corresponding high diversity at the causal 208 genetic level. Thus, it serves as a suitable model system to investigate the broader aspects of 209 genotype-phenotype relationships. Allelic composition at several glucosinolate biosynthetic 210 loci drive different glucosinolate profiles among A. thaliana ecotypes (Kliebenstein, 211 Kroymann, et al., 2001). These variations, however, are often in the form of presence and/or 212 expression of one or other copy of a duplicated gene, such as the AOP2 and AOP3 at the GS-213 ALK/GS-OHP locus (Kliebenstein, Lambrix, Reichelt, Gershenzon, & Mitchell-Olds, 2001), 214 or the MAM1/MAM2 at GS-ELONG (Kroymann et al., 2003), which complicates our 215 understanding of how genetic variations lead to metabolic properties of the enzymes encoded 216 by the respective genes. The Met-derived aliphatic GSLs are the most abundant form of 217 glucosinolates in A. thaliana and many Brassicaceae crops (Agerbirk & Olsen, 2012; 218 Benderoth et al., 2006; Halkier & Gershenzon, 2006; Kliebenstein, Kroymann, et al., 2001;

219 Kroymann et al., 2003; Textor et al., 2004; Textor, de Kraker, Hause, Gershenzon, & 220 Tokuhisa, 2007). The chain-elongation pathway of GSL biosynthesis is crucial for generating 221 the chain-length diversity of aliphatic GSLs and for connecting primary and specialised 222 metabolism. Although the evolution of core features of aliphatic GSL biosynthesis in 223 Arabidopsis has been studied (He et al., 2011; Sawada et al., 2009; Textor et al., 2004; 224 Wittstock et al., 2004), the molecular basis for diversity of function of MAM synthases and 225 the role of different MAM isoforms within A. thaliana accessions is not sufficiently 226 understood. The chain-length distribution of different aliphatic GSLs has so far been 227 attributed to the presence of different MAM isoforms, namely MAM1, MAM2 and MAM3 228 (Halkier & Gershenzon, 2006; Kroymann et al., 2003; Textor et al., 2004, 2007). By 229 investigating the differences in sequences of MAM synthases across A. thaliana ecotypes, we 230 show that the ecotypes can be broadly classified in two groups based on the similarity to 231 either of MAM1 and MAM2 genes as also expected from the genomic composition of the 232 GS_ELONG locus (cf. Figure 1). Correspondingly, the GSL profiles from different ecotypes can be broadly classified into two major groups based on the phenotypic distance $d_{i,i}^{P}$ 233 234 between different GSL profiles (cf. Figure 2). However, the groups classified based on either 235 of phenotypic distances or the similarity of MAM synthases are not identical. This points to a 236 more complicated relationship between the genotype and the associated GSL profiles. Thus, 237 estimating the pattern of GSL accumulation based solely on the distinction between MAM1 238 and MAM2 enzymes is not feasible.

239 Diversity of genes beyond classical QTLs

240 The heterogeneity in the genetic makeup of the metabolic genes across different A. thaliana 241 ecotypes and their associated metabolic phenotypes are an excellent tool for investigating the 242 mechanisms of adaptation and functional diversification (Mitchell-Olds & Schmitt, 2006; 243 Pigliucci, 2010). Comparing the diversity of genes of GSLs synthesis we expected to find the 244 highest diversity in genes of the chain-elongation and secondary modification, because these 245 steps contribute highly to the diversity of the GSLs. However, surprisingly, we found that the 246 level of diversity appears to be unrelated with the functional role of the gene within the GSL 247 metabolic pathway (cf. Figure 4). While as expected, the least diverse enzyme was SOT17, 248 part of the core synthesis, another enzyme of the core pathway, SUR1, was the fourth most 249 diverse from 30 enzymes (Figure 4). This is surprising, since loss of enzymes of the core 250 synthesis, such as UGT74B1 or SUR1 has a much higher impact on the total GSLs than loss 251 of enzymes of the side chain modification (Douglas Grubb et al., 2004; Keurentjes et al.,

252 2006; Mikkelsen, Naur, & Halkier, 2004). Thus, it seems that enzymes of all parts of GSL 253 biosynthesis can contribute to the diversity of the metabolites. Surprisingly, while 254 investigating the second least diverse enzyme BAT5, part of chain-elongation of aliphatic 255 GSLs, we found that it is highly diverse in the nucleotide sequence (see Figure 4B). It can be 256 a consequence of purifying natural selection that prevents the change of an amino acid 257 residue at a given position in a multiple alignment, thus favouring an excess of synonymous 258 versus non-synonymous substitutions. It is much more difficult, however, to explain the 259 results of *FMO-GSOX1*, which shows a large variation in amino acid sequence derived from 260 a relatively low variation in DNA sequence. Nevertheless, it is evident that FMO-GSOX1 261 favours non-synonymous substitutions versus the synonymous substitutions, possibly linked 262 to the function in the secondary modification of glucosinolates, responsible for large part of 263 the structural variation. Multiple genetic analyses revealed that, in general, few key QTLs 264 shape the metabolic phenotype. In contrast, our analysis detected diversity of the metabolic 265 genes across the whole pathway, irrespective of their association to a major QTL. However, 266 how far this variation in gene/protein sequence contributes to the phenotypic variation 267 remains to be elucidated.

268 The genotype-phenotype relationship

269 Nowhere is the contribution of subtle sequence diversity to variation in GSLs more apparent, 270 than in the MAM genes. In A. thaliana, the enzyme isoforms MAM1 and MAM2 catalyse the 271 formation of short-chain (3C and 4C) aliphatic GSLs, whereas the isoform MAM3 catalyses 272 the formation of both short-chain and long-chain (5C-8C) GSLs (Halkier2006). Indeed, 273 orthologues of MAM1 and MAM2 are also responsible for diversity of aliphatic GSLs across a 274 range of Brassica species (Kumar et al., 2019). The distinct function of the two genes was 275 confirmed by complementation of Arabidopsis mam1 mutant, when MAM1 from B. juncea 276 restored wild type GSL profile but MAM2 did not (Kumar et al., 2019). Our investigation of 277 genotypic and phenotypic distances between different Arabidopsis ecotypes showed that 278 some ecotypes have identical metabolic genotype but exhibit high diversity in their associated 279 metabolic phenotype, and vice-versa (cf. Figure 3). Therefore, the relationship between the 280 metabolic genotypes and the phenotypes are much more complicated than a link to one of the 281 MAM1/MAM2 gene pair. Indeed, the role of individual amino acid alterations between these 282 two genes demonstrates clearly that also SNPs can have a great effect on the phenotypes. 283 Thus, mutagenesis of serine to phenylalanine at position 102 and alanine to threonine at 290, 284 parts of active region of MAM1 changed the distribution of C3 to C4 GSLs (Kroymann et al.,

285 2001) in A. thaliana. Alterations in four other amino acids in B. juncea MAM1 affected the 286 kinetic properties of the enzyme to more MAM2-like and vice versa (Kumar et al., 2019). 287 Also, in Arabidopsis MAM1 alteration of further three amino acids resulted in changes of the 288 pattern of elongation products in vitro (Petersen et al., 2019). To further investigate the 289 impact of MAM synthases on chain-length distribution of aliphatic GSLs, we analysed the 290 polymorphisms in the active region of MAM synthases. From our analysis of the diversity of 291 the active region of MAM synthases, we conclude that MAM1 is highly variable across its 292 active region and accumulate up to 13 polymorphic amino-acid residues at 17 different 293 locations in the active region. Whereas, the active region of MAM3 is comparatively 294 conserved and only accumulates a maximum of 4 polymorphic residues at 3 locations in the 295 active region, naturally (cf. Figure 5). It is known that polymorphisms in active site of MAM 296 synthases change the specificity of a metabolic enzyme towards respective substrates of 297 aliphatic GSL biosynthesis (Kumar et al., 2019; Petersen et al., 2019). This results in 298 different composition and the total accumulation of aliphatic GSLs across different 299 Brassicaceae species, including Arabidopsis thaliana (Kroymann et al., 2001; Kumar et al., 300 2019; Petersen et al., 2019). The above analysis, however, only showcases one of the two 301 possibilities by which a genotype can exhibit a metabolic phenotype. The gene regulatory 302 networks can also change the expression of metabolic genes, which in turn changes the 303 enzyme abundance and thus results in different metabolic phenotypes (de Kraker & 304 Gershenzon, 2011; Kumar et al., 2019; Petersen et al., 2019). Although, numerous studies 305 have shown that a multitude of genes and underlying regulatory processes are involved in the 306 diversity of specialised metabolites such as glucosinolates (Chan et al., 2010; Koornneef, 307 Alonso-Blanco, & Vreugdenhil, 2004; Kumar et al., 2019; LASKY et al., 2012; Petersen et 308 al., 2019), interpreting the findings in the context of metabolic properties is highly 309 challenging. This is particularly due to a missing stringent definition of the genotype-310 phenotype relationship, which can hardly be expected to be derivable from a single 311 methodology but rather requires a comprehensive platform of combined experimental and 312 theoretical strategies (DIZ, MARTÍNEZ-FERNÁNDEZ, & ROLÁN-ALVAREZ, 2012; 313 Sharma, 2018; Weckwerth, Wenzel, & Fiehn, 2004).

314 Conclusion

Altogether we show here that the control over phenotypic diversity in glucosinolates is potentially spread over the whole pathway. On the example of MAM1 and MAM2, responsible for side chain elongation of Met-derived glucosinolates, we revealed that 318 sequence variation beyond the presence of one or the other isoform contributes to the 319 variation in chain length. The present study thus points to the necessity to pay attention to 320 variation beyond the classical ON/OFF features of key metabolic QTLs, for investigating the 321 diversity of specialised metabolic pathways, such as glucosinolates. Since the recent efforts 322 towards unravelling the genotype-phenotype relationships focus at either experimental 323 studies with a selection of genotypes or computational approaches to correlate the observed 324 experimental observations, it is crucial to develop frameworks that integrate multi-omics data 325 with fundamental rules of metabolic modelling to fully understand how particular genotype is 326 reflected in a phenotype.

327 Materials and Methods

328 Genotypic data

329 Information about the nucleotide and amino acid composition of 30 GSL biosynthesis genes from 72 A. thaliana ecotypes was taken from the 1001 genomes project (Jorge et al., 2016). 330 331 The reason behind selecting these 72 ecotypes was the availability of Met-derived GSL 332 composition under identical environmental conditions. To obtain the gene sequences of the 333 ecotypes of interest, we used an inhouse R-script that converts the TAIR10 version of SNP 334 (single nucleotide polymorphisms) files provided by the 1001 genome database into an R-335 object. This R-object is a sparse matrix containing the nucleotide information for each 336 ecotype at each locus in the reference genome coded as numbers. Non-polymorphic sites are 337 coded as 0, polymorphic ones as 1,2,3,4 or 5 depending if at the specific locus an A, C, G, T 338 or indel was observed. From this R-Object we could extract the nucleotide sequences of a 339 specific ecotype for each coding region of interest. To obtain the amino acid sequence we 340 used the function 'translate' from the R-package 'seqinr'.

341 Phenotypic data

Experimental data of Met-GSLs concentrations were obtained from Chan et al. (2010) and Kliebenstein et al. (2001). The final set of GSL data is given in Table 1. The data is composed of normalised concentrations of six aliphatic GSLs, referred to as 3C to 8C, from 72 different ecotypes of *A. thaliana* under controlled and identical experimental conditions.

346

347 Calculation of diversity using Shannon entropy

348 The nucleotide/amino-acid composition of the coding regions of GSL genes from 72 A.

349 *thaliana* ecotypes is described as a set of relative probability, $p_{i,i}$, for the i^{th} nucleotide/amino-

acid (i = 1, 2, ..., n) in the j^{th} ecotype (j = 1, 2, ..., 72). Then, the diversity of each position

in the coding region can be quantified by Shannon entropy (Shannon, 1948),

$$H_i = -\sum_j^n p_{i,j} \log p_{i,j}$$

352 H_i will vary from zero, when the i^{th} nucleotide/amino-acid is same across all 72 ecotypes, to 1

353 when the probability is equal for observing all nucleotides/amino-acids at same locus across

354 72 ecotypes. Moreover, to get an estimate of diversity of a nucleotide/amino-acid sequence of

length *n*, we calculate the average entropy H^{avg} as

$$H^{avg} = \frac{1}{n} \sum_{i}^{n} H_i$$

356 Phylogenetic tree reconstruction

Amino acid sequences of the MAM loci from the 72 ecotypes were aligned using 'mafft' ver. v7.407 (Katoh & Standley, 2013) with the parameter '--maxiterate 1000 --globalpair -phylipout' to obtain a multiple sequences alignment in phylip format. This was then used as input for phyml (20120412) (Guindon et al., 2010) to reconstruct the maximum likelihood phylogenetic tree using the LG substitution model. The tree was visualised with FigTree v1.4.2 (http://tree.bio.ed.ac.uk/software/figtree/).

363 Resources

The data and Python scripts used to produce the results presented in this manuscript are available with instructions at (<u>https://gitlab.com/surajsept/GTvsPT</u>).

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370 Author contributions

- 371 SS, OE and SK planned and designed the research. SS performed the computational work and
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543

Figures

Figure 1: Mid-point rooted phylogenetic tree showing the evolutionary relationship among coding regions of annotated

MAM1 gene of 72 Arabidopsis thaliana ecotypes. The red branch represents ecotypes showing higher similarity to the

MAM2 sequence. The scale bar is substitutions per position.



- 552 553 554 Figure 2: Clustered heatmap of the GSL composition across 72 different A. thaliana ecotypes. The top cluster is composed of
- ecotypes having high concentration of 3C GSLs, the lower cluster corresponds to the ecotypes accumulating high

concentrations of 4C GSLs.



555

557 Figure 3: The genotypic versus phenotypic distance. Every ecotype is assigned to possess either MAM1 or MAM2, based on

the sequence similarity of annotated MAM1 gene. Each dot represents a pair of ecotypes. The colours red and blue denote

558 559 pairs, where both ecotypes show high similarity to the coding region of MAM1 and MAM2 sequence, respectively, the green 560 dots denote heterogeneous pairs.



563 Figure 4: Diversity of GSL genes from 72 A. thaliana ecotypes. (A) The bars represent the diversity of the amino-acid (AA)

residues of respective GSL genes. The bars are colour coded to denote genes from the chain-elongation process, core-

structure formation and secondary chain modifications by red, blue and green colours, respectively. (B) The bars represent
 the diversity of the nucleotides (NT) of respective genes. The bars are colour coded as in (A). (C) Diversity of NT residues

567 plotted against the diversity of AA residues of respective GSL genes. Each dot is colour coded to denote genes as in (A) and

568 (B). The black dashed line is a linear regression line.



Figure 5: Polymorphisms in the active region of MAM synthases. (A) Substitutions in the active sites of MAM1 versus the
 Genotypic distances. (B) Substitutions in the active sites of MAM1 versus the Genotypic distances. (C) Polymorphisms in the
 active region of MAM1 and MAM3.



Supplementary Figures: 580

581 Figure 1: Diversity of GSL enzymes from different ecotype subgroups. The bars represent the diversity of the amino-acid

582 residues of respective GSL genes. The blue bars denote all 72 ecotypes, while orange and green denote ecotypes having 583 MAM1 and MAM2, respectively.



585

586 Figure 2: Diversity of GSL enzymes from different subgroups of ecotypes. The bars represent the diversity of nucleotide 587 sequences of respective GSL genes. Colour coding is same as Supplementary Figure 1.

