Laminarin-triggered defence responses are geographically dependent for natural populations of *Solanum chilense*.

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10 Short title: diversity in laminarin responses in wild tomato

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48 Highlight

49 Large-scale screenings reveal geographically distinct intraspecific differences in the 50 dominant physiological pathogen defence responses upon glucan elicitor treatment in a wild 51 tomato species.

52

53 Abstract

54 Natural plant populations are polymorphic and show intraspecific variation in resistance 55 properties against pathogens. The activation of the underlying defence responses can depend 56 on variation in perception of pathogen-associated molecular patterns or elicitors. To dissect 57 such variation, we evaluated the responses induced by laminarin, (a glucan, representing an 58 elicitor from oomycetes) in the wild tomato species *Solanum chilense* and correlated this to 59 observed infection frequencies of *Phytophthora infestans*.

60 We measured reactive oxygen species burst and levels of diverse phytohormones upon 61 elicitation in 83 plants originating from nine populations. We found high diversity in basal 62 and elicitor-induced levels of each component. Further we generated linear models to explain 63 the observed infection frequency of *P. infestans*. The effect of individual components differed 64 dependent on the geographical origin of the plants. We found that the resistance in the 65 southern coastal region, but not in the other regions is directly correlated to ethylene 66 responses and confirmed this positive correlation using ethylene inhibition assays.

67 Our findings reveal high diversity in the strength of defence responses within a species and 68 the involvement of different components with a quantitatively different contribution of 69 individual components to resistance in geographically separated populations of a wild plant 70 species.

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72 Keywords: Diversity; Early Immune Response; Ethylene; Laminarin; Phytohormones;
73 *Phytopthora infestans*; Reactive Oxygen Species; Resistance; *Solanum chilense*; Tomato

74 Introduction

75 Plant defence responses against pathogens in natural populations are often polymorphic 76 (Kahlon and Stam, 2021a). The resistance of the host can result from the major resistance 77 proteins or from polygenic defence mechanisms against the pathogen (Vanderplank, 1963). 78 The later type of resistance mechanism is often considered as a basal defence, whereas major 79 gene-mediated resistance is observed as pathogen genotype dependent. Genes encoding 80 nucleotide-binding domain leucine-rich repeat-containing proteins (NLR) are one example of 81 major resistance genes. NLR genes have been studied in natural plant populations and are 82 reported to be diverse at the genetic level (Stam et al., 2019a, Van de Weyer et al., 2019; 83 Witek et al., 2021). Similarly, members of the receptor-like proteins (RLPs) family show 84 intraspecific variation in the presence-absence of defence responses or variability in 85 expression patterns of these genes (Van der Hoorn et al., 2001; Kruijt et al., 2005; Kahlon et 86 al., 2020, Steidele and Stam, 2020). In many cases, major gene mediated resistance is 87 complete. By contrast, basal resistance is defined as quantitative and pathogen race-non-88 specific (Vanderplank, 1963). This might partially be explained by its polygenic nature, and 89 because underlying defence reactions can be activated upon exposure to elicitors or 90 conserved pathogen-associated molecular patterns (PAMPs). Flagellin peptides (flg22, flgII-91 28) and chitin have been the dominant PAMPs for studies on resistance mechanisms in plants 92 against pathogens from bacterial and fungal lineages respectively. Many additional PAMPs 93 have been identified e.g.VmE02 homologs, produced by various fungi and oomvcetes, can 94 trigger immunity response in N. benthamiana (Nie et al., 2018) and peptide elicitor fractions 95 from several Fusarium spp. activate basal defence mechanisms in Arabidopsis thaliana 96 (Coleman et al., 2021). One other example is laminarin, which is perceived in different plant 97 species like Nicotiana tabacum (Klarzynski et al., 2000; Ménard et al., 2004), Grapevine, 98 Vitis vinifera (Aziz et al., 2003), A. thaliana (Ménard et al., 2004), tea, Camellia sinensis 99 (Xin et al., 2019), Nicotiana benthamiana, Hordeum vulgare, Brachypodium distachyon 100 (Wanke et al., 2020) and Olive, Olea europaea (Tziros et al., 2021). Laminarin is an 101 oligometric β -1,3-glucan with β -1,6-glucan branches. β -1,3 and β -1,6-glucan are the major 102 components of the oomvcete cell wall (Aronson et al., 1967) and may induce defence 103 responses similar to those provoked by elicitors from the oomycete lineage.

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105 Several molecular mechanisms have been shown to play an important role in basal defence 106 responses. Many of these responses happen shortly upon contact with the pathogen. In some

107 plant-pathogen interactions basal immune responses can be quantified within the first minutes 108 of the interaction with the pathogen or pathogen-specific molecules by measuring reactive 109 oxygen species (ROS) production (Torres et al., 2006). The fine-tuning in the production of 110 ROS is one important cue toward activating resistance mechanisms which lead to the 111 production of various phytohormones or activation of downstream defence regulators 112 (Ramirez-Prado et al., 2018). Roberts et al. (2019) showed the amount of ROS production 113 varied when different tomato (Solanum lycopersicum) accessions were treated with flagellin 114 peptides (flg22, flgII-28). Besides the production of ROS, phytohormones present or induced 115 in the plant can greatly influence the resistance outcome. A higher level of salicylic acid (SA) 116 is important in activating defence response in cultivated tomato leaves against P. infestans 117 (Jeun et al., 2000). Genes involved in ethylene (ET) and SA pathways are important in N. 118 benthiamana after infection with P. infestans (Takemoto et al., 2018). A study on potato 119 shows that upon infection with P. infestans large sets of genes are upregulated at multiple 120 time points post-inoculation. These included key marker genes involved in the jasmonic acid 121 (JA) acid signalling pathway and genes involved in primary and secondary metabolite 122 pathways (Tian et al., 2006). In cultivated tomato the negative role of abscisic acid (ABA) in 123 resistance against *Botrytis cinerea* is regulated by repressing SA signalling (Audenaert *et al.*, 124 2002). Whereas resistance against Alternaria solani is enhanced upon exogenous ABA 125 application through defence-related gene activation and defense-related enzymatic activity of 126 phenylalanine ammonia-lyase (PAL), polyphenol oxidase (POD) and peroxidase (PPO) 127 (Song et al., 2011). Exogenous application of indoleacetic acid (IAA) in the soil resulted in 128 Fusarium oxysporum lycopersici disease suppression in tomato plants (Sharaf and Farang, 129 2004).

130 The different components of the phytohormone signalling pathways can have positive or 131 negative feedback effects on each other and thus form a complex interactive signalling 132 network (Pieterse *et al.*, 2009). These complex network topologies generated an hypothesis 133 that one or multiple components involved in the resistance need to pass a certain threshold, in 134 order for defence to be functional (Windram and Denbi 2015). Which factors are dominant 135 might differ dependent on the origin of a plant or population and the pathogen in question.

136 Such differences in dominant effective defence responses in different *A. thaliana* accessions 137 are shown by Velásquez *et al.* (2017) against the bacterial pathogen *Pseudomonas syringae* 138 pv. tomato (*Pst*) DC3000. They found that *Pseudomonas syringae* pv. tomato (*Pst*) DC3000 139 resistance was shown to be mainly mediated by an increased level of the phytohormone 140 salicylic acid (SA) in three accessions, whereas other mechanisms were dominant in the other

141 resistant accessions. Another study showed that between six *A. thaliana* accessions large 142 variation in JA and SA- associated basal resistance resulted in varied resistance against the 143 nectrotrophic pathogen *Plectosphaerella cucumerine* and the hemibiotrophic bacterium *P.* 144 *syringae* respectively (Ahmad *et al.*, 2011).

145 We used a wild tomato species Solanum chilense to elucidate molecular cues behind the 146 diversity in resistance against the oomycete, P. infestans (Stam et al., 2017; Kahlon et al., 147 2021). S. chilense is a suitable organism to study the variation of molecular responses 148 associated with basal defence mechanisms. Populations of S. chilense are geographically 149 structured in four distinct groups based on genomic studies (Böndel et al., 2015; Stam et al., 150 2019a). The two southern groups are recent expansions of the species, are genetically more 151 divergent and might be developing into new subspecies (Raduski and Igic, 2021). Thus the 152 system provides a strong genetic structure. Previously, we found variation in defence 153 responses against the apoplastic fungal leaf pathogen Cladosporium fulvum (syn. Fulvia 154 fulva, Passalora fulva), with complete loss of pathogenic protein recognition in plants from 155 the southern groups (Kahlon et al., 2020). In order to assess phenotypic variation in 156 resistance, we also quantified the number of successful infection events in S. chilense plants 157 after drop inoculation with various other leaf pathogens. We showed clear differences in the 158 frequency of successful infections after inoculation of three common filamentous pathogens 159 in different S. chilense populations (Stam et al., 2017). In a recent report (Kahlon et al., 2021) 160 we showed that differences in *P. infestans* resistance between the geographically distinct 161 populations of S. chilense are predominantly driven by the host genotype and can likely be 162 attributed to differences in basal resistance, rather than isolate-specific resistance.

163 Here, we aim to dissect the various possible immune responses in *S. chilense* populations. We 164 specifically investigate the early basal defence responses in *S. chilense* upon challenge with 165 the nonspecific elicitor laminarin. We confirm that laminarin elicits a subset of defence 166 responses triggered by *P. infestans*, we report high diversity in several key regulators of basal 167 immune responses in *S. chilense* within and between populations of the species within the 168 first hours of infection, and we assess their individual and joint effect on the interaction 169 outcome, by comparing these results to our previously published infection data (Kahlon *et al.*, 170 2021).

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172 Materials and methods

173 Plants material used and maintained

We used 83 plants of *S. chilense* originating from nine populations (accessions) (8-10 plants 175 each): LA1958, LA1963, LA2747, LA2932, LA3111, LA3786, LA4107, LA4117A and 176 LA4330. The seeds of these populations were procured from the C. M. Rick Tomato Genetics 177 Resource Center of the University of California, Davis (TGRC UC-Davis, 178 <u>http://tgrc.ucdavis.edu/)</u> where the populations were originally collected as random collection 179 of seeds from the wild populations and are now maintained and propagated at the TGRC to 180 maintain genetic diversity. Procured seeds were sown and plants were maintained in 181 controlled greenhouse conditions (16h light and 24^oC temperature at daytime and 18-20^oC at 182 night) at TUM's plant technology center. Each plant used in this study was at least a year old. 183 Plants were maintained throughout the experiments by cutting them every two weeks. Each 184 population used in this study represents one of four geographical locations of the species 185 habitat and were originally collected during different years from wild populations. Each 186 individual plant within a population is genetically unique.

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188 Evaluation of laminarin potential to activate early immune responses similar to *P*. 189 *infestans* using 3' RNAseq

We selected the central population LA3111 to evaluate differentially expressed genes in the transcriptome upon challenging with *P. infestans* Pi100 (3000 sporongia/ml) and laminarin (1mg/ml, Sigma-Aldrich) treatment (using spray inoculation). We use LA3111 population because the reference genome of *S. chilense* was generated from an individual from this population (Stam *et al.*, 2019b). To measure the general defence responses in the population, the experiment was done on nine plants and all plants were pooled per treatment for the RNA extraction. Detached leaves were kept upside down in plastic boxes containing wet tissue beds and treated with water, laminarin or *P. infestans*. The boxes were kept at 18-20°C for 3 hours and samples were taken and snap-frozen in liquid nitrogen. RNA was isolated using Qiagen[®] RNeasy plant mini kit according to the instruction manual. Each treatment consisted of pooled samples of nine plants and four technical replicates of each treatment.

201 3'RNA libraries were prepared according to the manufacturer's protocol using the QuantSeq
202 3'mRNA-Seq Library Prep Kit (Lexogen, Vienna, Austria). Sequencing was performed on a
203 HiSeq2500 (Illumina, San Diego, CA, USA) with single-end 100bp reads using Rapid SBS
204 v2 chemistry. The raw sequencing reads in FastQ format were trimmed to remove adapter
205 sequences using Trimmomatic v0.39 (Bolger *et al.*, 2014). The reads were quality filtered and
206 trimmed using the following settings: LEADING:3, SLIDINGWINDOW:4:15, MINLEN:40.
207 HISAT2 (Kim *et al.*, 2015) was used to align sequencing reads to a reference genome (Stam

et al., 2019b). After alignment, featureCounts (Liao *et al.*, 2014) was used to identify the number of reads that mapped to genes. For feature Counts, all entries tagged as 'gene' were extracted from the gff annotation files and by adjusting the gene_id and transcript_id identifiers, this gene list was converted into a gtf annotation file. The downstream region of every gene was extended by 1kb (the extension stops when it hits the next gene start site). FeatureCounts was modified to search for the tag 'gene' instead of the default 'exon' tag.

214 Differential gene expression analysis was carried out using the R package DESeq2 (Love *et* 215 *al.*, 2014). DESeq2 uses the output of featureCounts to estimate the fold change in gene 216 expression between different treatment groups. Default parameters from the DESeq2 package 217 were applied and differentially expressed genes (DEGs) showing adjusted *p*-value<0.05 were 218 considered significant.

219 Gene Ontology (GO) enrichment analysis was based on previously annotated ontologies 220 (Stam *et al.*, 2019b). GO terms were selected for all candidate genes. The Background 221 frequency of each GO term is the number of genes annotated to this GO term in all genes, 222 while sample frequency is the number of genes annotated to that GO term in the list of DEGs 223 in this sample.

For the manual inspection of the functions of the differentially expressed overlapping gene candidates in laminarin and *P. infestans* treated samples, we performed a BLAST search, extracted gene names and functional annotation from the best hits and if needed, performed a literature search for papers that described the functions of the described gene candidates. (See 10.5281/zenodo.5101308)

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230 Gene expression analysis of the key indicators of phytohormones pathways via qPCR

To independently evaluate phytohormone regulation in response to laminarin (1mg/ml) elicitation we tested the expression level of key indicators of three well-known defence and phytohormones *1-aminocyclopropane-1-carboxylic acid synthases 2*, *ACS2* from the ET and pathway (Gravino *et al.*, 2015), *isochorismate synthase*, *ICS* (Di *et al.*, 2017) and *phenylalanine ammonia-lyase*, *PAL* (Peng *et al.*, 2004) from the SA pathway and *lipoxygenase D*, *LOXD*; (Heitz *et al.*, 1997) from the JA pathway, in individual plant leaf discs of *S. chilense* upon treatment with laminarin and compared it to mock-treated (water) and leaf discs. *S. chilense* reference names of the genes are provided in Table S1.

239 Leaf discs of the plant LA1963-02 (chosen due to its high resistance observed in Kahlon *et* 240 *al.*, 2021) were treated with laminarin and MilliQ-H₂0 treated leaf discs served as control. 241 Experiments were performed on three different dates in three independent replicates each.

242 Samples were treated for 1.5 hours, snap-frozen in liquid nitrogen and ground to a fine 243 powder with a mortar and pestle. RNA was extracted using the Qiagen[®] RNeasy plant mini 244 kit according to the instruction manual. cDNA synthesis was performed using a Qiagen 245 QunatiTect[®] reverse transcriptase kit according to the instruction manual. Quantitative PCR 246 (qPCR) was performed on the synthesised cDNA using Takyon[™] Low ROX SYBR[®] master 247 mix ddTTP blue (Eurogentec Liège, Belgium). qPCR was performed in three technical 248 replicates and a non-template control was included. Primer pairs for each tested gene are 249 indicated in Table S1, TIP-41 (Fisher et al., 2013; Nosenko et al., 2016) was used as a 250 housekeeping gene for normalization and primer efficiency was performed for all the primer 251 pairs and are shown in Table S1. The PCR reaction comprised of 10µl SYBR Green-ROX 252 Mix, 0.3µM of forward primer and 0.3µM of reverse primer, 3µl of cDNA and volume was 253 adjusted to 20µl with MilliQ-H₂0. The thermal cycling profile was set to a hot start at 95°C 254 for 3 minutes, followed by 40 cycles of amplification (95°C for 30 seconds, 60°C for 30 255 seconds, 72°C for 1 minute), 1 cycle melting (95°C for 30 seconds, 65°C for 30 seconds, 256 95°C for 30 seconds), and in the end 1 cycle at 72°C for 10 minutes. Melting curve 257 temperatures were recorded at the end of the cycle for quality control. Data were evaluated 258 with the software Agilent Aria MX 1.7 and relative gene expression was calculated based on 259 the Livak and Schmittgen method (2001).

260 ROS production measurement

To measure ROS production upon elicitation with laminarin, we performed a 96-well plate assay based on chemiluminescence as described by Kahlon and Stam (2021b). Leaf discs from leaves of mature plants were made with a biopsy punch (4mm, KAI Medical Solingen, Germany) and incubated in white 96-well flat-bottom plate overnight in 200µl of 20mM MOPS (pH 7.5) at room temperature. The next day buffer was removed and wells were supplemented with 75µl HRP mix (10µM horseradish peroxidase (HRP) and 10µM L012). A baseline reading was performed for the initial 10 minutes using a Luminoskan Ascent (Thermo Scientific) and then laminarin (1mg/ml final concentration) dissolved in MOPS was added to the plate at 4 wells/plant. For each population, the assay was performed on three different dates with four technical replicates each for treatment and mock (MOPS) per date. In addition, we performed ROS production measurements with flg22 at a final concentration of 500nM in population LA4330 (7 plants) to compare it with specificity of laminarin in ROS production. Normalization of data was performed by first averaging over the initial 6-10 and minutes baseline reading, followed by normalization to the mock treatment for the treated

275 leaf discs.

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277 Phytohormone measurements: ET measurements

278 Leaf discs were obtained with a 4mm biopsy punch and incubated overnight in Petri dishes 279 containing milliQ-H₂O at room temperature. Following overnight incubation, three leaf discs 280 were added to glass vials (5ml) containing 300µl of milliQ-H₂O. Laminarin was added in a 281 final concentration of 1mg/ml to 3 glass vials (samples) containing three leaf discs of one 282 plant and milliQ-H₂O in three separate glass vials containing leaf discs for same plant served 283 as a negative control. Upon addition of elicitor or water, the glass vials were sealed with septa 284 (Carl Roth GmbH). Samples were incubated for three hours at a shaker at ~ 20-50 rpm 285 (Heidolph Polymax 2040). 3 hours post-incubation, 1ml of air was retrieved from each 286 samples with a syringe through the rubber cap and injected into a Varian 3300 gas 287 chromatography machine containing AlO₃ column with length 1m and 225°C detector 288 temperature and 80°C column and injector temperature. The gases used for the separation of 289 ET from the sample were H₂, N₂ and O₂ at 0.5 MPa each. The amount of ET was calculated 290 based on the standard calculation as developed by Von Kruedener *et al.* (1994) using the area 291 under the curve (AUC). In total, we measured up to nine samples per plant on three different 292 dates, each date containing up to three samples.

293

294 Phytohormone and their derivatives measurements: Measurement of SA, JA, ABA, 295 IAA, phaseic acid (PA) and dihydrophaseic acid (DPA):

Samples for measurements of these six compounds were also prepared based on the leaf disc treatment method. 150-200 leaf discs were made per plant using a 4mm diameter biopsy punch and incubated overnight in Petri dishes containing milliQ-H₂0. The next day for each plant a 6-well plate filled with milliQ-H₂0 was prepared for elicitation containing 25-30 leaf discs per well. Three wells were elicited with laminarin (1mg/ml) and in the remaining three wells milliQ-H₂0 was added as a control. The plates were incubated for three hours at a shaker at ~ 20-50 rpm. Following the treatments, the leaf discs were transferred to 2ml Eppendorf tubes and residual water was pipetted out before snap-freezing the samples in 304 liquid nitrogen.

305 Fine powder from the plant material was obtained after grinding the frozen leaf discs with 306 mortar and pestle in liquid nitrogen. The samples were then processed for extraction of the 307 phytohormones and their derivatives as described by Chaudhary *et al.* (2020), with minor 308 modifications. 50-200mg ground material was transferred to a 2ml bead beater tube (CKMix-

309 2ml, Bertin Technologies, Montigny-le-Bretonneux, France). 20µl of internal standard 310 solution containing indoleacetic acid-d2 (Sigma Aldrich, Steinheim, Germany) (2.5µg/ml), 311 salicylic acid-d4 (Olchemim, Olomouc, Czech Republic) (2.5µg/ml), (+) cis, trans-abscisic 312 acid-d6 (Sigma Aldrich, Steinheim, Germany) (2.5µg/ml), and (-) trans-jasmonic acid-d5 313 (25µg/ml) (Santa Cruz, Dallas, TX, USA) were dissolved in acetonitrile and added to the 314 samples and incubated for 30 minutes at room temperature. Following that 1ml of ice-cold 315 ethyl acetate (Art. 864, Merck, Darmstadt, Germany) was added to the samples and stored 316 overnight at -20°C. The next day samples were shaken for 3X20 seconds using the bead 317 beater (Precellys Homogenizer, Bertin Technologies, Montigny-le-Bretonneux, France) at 318 6000rpm with 40 seconds breaks in-between. The material was then filtered with a 0.45µm 319 pore size filter (Sartorius, Darmstadt, Germany) using a Minisart® syringe. The filtrate was 320 transferred to 2ml tubes and vacuum dried. Then samples were reconstituted in 70µl of 321 acetonitrile and sonicated for 3 minutes. 2µl of the sample from the HPLC tubes (glass vials) 322 were injected into the LC-MS/MS system. The MS method used measured positive and 323 negative ionization mode within one run (polarity switching). Negative ions were detected at 324 an ion spray voltage of -4500 V (ESI-) using ion source parameters: curtain gas (35 psi), 325 temperature (550°C), gas 1 (55 psi), gas 2 (65 psi), collision activated dissociation (-3 V), 326 and entrance potential (-10 V). Positive ions were detected at an ion spray voltage at 4500 V 327 (ESI+) using ion source parameters: curtain gas (35 psi), temperature (550°C), gas 1 (55 psi), 328 gas 2 (65 psi), collision activated dissociation (-3 V) and entrance potential (10 V) and 40°C 329 column oven temperature was at a QTRAP 6500+ mass spectrometer (Sciex, Darmstadt, 330 Germany). MS/MS fragmentation was obtained and samples were separated by ExionLC 331 UHPLC (Shimadzu Europa GmbH, Duisburg, Germany) using 100 × 2.1 mm2, 100 Å, 1.7 332 µm, Kinetex F5 column (Phenomenex, Aschaffenburg, Germany). Solvent used for 333 separation were (A) 0.1% formic acid in water (v/v) and (B) 0.1% formic acid in acetonitrile 334 (v/v) with a flow rate of 0.4 ml/minute. Chromatographic separation was performed with the 335 gradient of 0% B for 2 minutes, increased in 1 minute to 30% B and in 12 minutes to 30% B, 336 increased in 0.5 minute to 100% B, held 2 minutes isocratically at 100% B, decreased in 0.5 337 minute to 0% B, and held 3 minutes at 0% B. Phytohormone quantification was performed 338 based on comparison with standard curves prepared with purified hormones and using AUC. 339 The final concentrations were obtained in nanograms of hormone per gram of fresh weight of 340 the sample.

341

342 Infection data

343 Data on *P. infestans* infections were taken from Kahlon *et al.* (2021), using the same methods 344 that were previously described in Stam *et al.* (2017). In these studies, detached leaves were 345 drop infected with a *P. infestans* solution (3000 sprongia/ml). All leaflets of the compound *S.* 346 *chilense* leaves were infected with a single drop and the infection frequency (IF) was 347 calculated per leaf and summarized per plant an population. The data originate from the exact 348 same plants as those used in this study.

349

350 Statistical analysis of the data, Pearson's correlation and linear mixed models for 351 infection frequencies with components of basal immunity and stress-related 352 phytohormones

353 All the data analyses were performed in the R software (version 3.4.4, R core Team, 2020). 354 ANOVA was performed with the function aov(), and post hoc Tukey tests with the function 355 TukeyHSD(), from the package {stats}. When the *p*-value was lower than 0.05 it was 356 considered significant. Pearson's correlation was performed using the function cor(). The 357 analyses were done for the 83 plants for which IF scores were available (Kahlon *et al.*, 2021). 358 Figures were made using the R package {ggplot2}.

359

360 Validation of ET accumulation in delivering resistance in individuals from the southern 361 coastal population

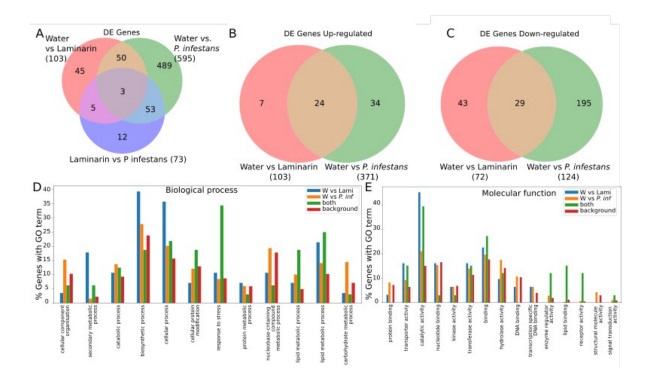
362 ET validation experiments were performed on two individuals (plant 05 and plant 10) from a 363 southern coast population, LA4107, selected based on Pearson's correlation among ET 364 production and infection frequency. ET measurement in the leaf discs was performed as 365 described above. ET blocking was performed by adding 5μ M aminoethoxyvinyl gylcine 366 (AVG) (Sigma) to the samples and 3 hours post-treatment ET was measured by gas 367 chromatography. The infection frequency upon treatment with 5μ M AVG was determined 368 using detached leaf infection assay as described in Kahlon *et al.* (2021): detached leaves were 369 surface sterilized with 70% ethanol and kept upside down (adaxial side facing upwards) in 370 plastic boxes containing a wet tissue bed. The leaf set for AVG treatment was kept on a wet 371 tissue bed made with water containing AVG (5μ M) and leaves were sprayed with 5μ M AVG 372 following drop inoculation with *P. infestans* isolate Pi100 (3000 sporangia/ml). The 373 experiment was repeated on eight individual leaves per treatment. Throughout the 374 experiment, 18-20°C temperature was maintained and boxes were kept in dark. The infection 375 outcome was taken at 7 days post-inoculation.

377 **Results**

378 Laminarin elicits a transcriptional response overlapping with that induced by *P*. 379 *infestans*

380 First, we set out to confirm whether the purified glucan elicitor laminarin can elicit 381 oomycete-like early defence responses in *S. chilense*. Therefore, we infected *S. chilense* 382 plants of population LA3111 with *P. infestans* or treated the plants with laminarin and 383 measured the transcriptional response after 3 hours.

384 As expected, infection with P. infestans triggered strong transcriptional responses. In total, 385 we measured 595 differentially expressed genes (false discovery rate adjusted *p*-value<0.05) 386 upon P. infestans infection (371 up-regulated and 224 down-regulated). Laminarin treatment 387 results in 102 differentially regulated genes (31 up-regulated and 72 down-regulated). (Table 388 S2). More than 50% of the genes that were differentially expressed after laminarin treatment 389 were overlapping with the P. infestans-associated response. For the upregulated gene 390 fraction, the overlap was 77%. Only a small number of DEGs (12) can be uniquely detected 391 in a direct pairwise comparison between the Laminarin-treated and Phytophthora-treated 392 samples, thus the false positive discovery rate in this experiment is likely lower than 2%. 393 (Figure 1A-C). To validate our hypothesis that laminarin triggers a decomplexified defence 394 response, we analysed the annotations of the overlapping gene lists. 35% of the overlapping 395 genes are associated with the Biological process: Stress Response and the overlapping 396 fraction is significantly more often annotated with the GO terms enzyme regulator activity 397 and receptor activity (chi-square test, p < 0.01, Figure 1D-E, Table S3-S4). Moreover, 398 homologs of more than half of the overlapping genes are reported in the literature to be 399 involved in defence responses (Table S5). We found homologs of regulators of the plant ROS 400 response (SOLCI005830700, Peroxidase CEVII), key regulators of defence hormone 401 signalling like ER5 (SOLCI004643500, ET response); LOX1 (SOLCI003764800, JA 402 signalling) or PAL3 (SOLCI000597200, SA signalling) and upregulation of Mitogen-403 activated protein kinases (MAPKs, SOLCI002491100). This supported that laminarin can 404 trigger a subset of oomycete-associated defence responses and is a suitable compound to 405 study variation in basal defence responses in S. chilense.



407 Figure 1: RNAseq analysis of the central population LA3111 (nine plants pooled per treatment) of *S.* 408 *chilense* 3 hours after *P. infestans* (3000 sporongia/ml), laminarin (1mg/ml) and water treatment. 409 Differentially expressed genes (DEGs) overlap in different treatment: overall (a), up-regulated (b) and 410 down-regulated (c), Gene Ontology (GO) analysis of RNA-seq data showing % genes with signals for 411 gene ontology terms for biological and molecular function for laminarin Vs *P.infestans* treatment 412 (d,e).

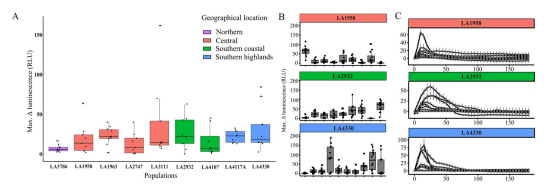
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414 To independently verify the involvement of laminarin-specific phytohormone-associated 415 defence responses, we evaluated the expression of key regulators described in literature of 416 three phytohormones, ET, SA and JA in an *S. chilense* individual from a different central 417 population: LA1963 (plant 02). We observed an up to 7.8-fold increase in expression of *S.* 418 *chilense ACS2* (SOLCI000989600), a key regular in the ET pathway in laminarin treated 419 samples as compared to water-treated samples (Figure S1). Next, we looked into key 420 regulators from the two known pathways for SA regulation. We observed that *PAL*-like 421 transcripts (SOLCI002546900) showed up to 14-fold increase in laminarin-treated samples as 422 compared to water-treated controls. *ICS* (SOLCI004470400), the key regulator of the second 423 SA pathway was downregulated (Figure S1). For the JA pathway, we performed qPCR on the 424 *LOXD* (SOLCI003768300) gene and observed an up to 10-fold increase in the laminarin 425 treated samples (Figure S1). Thus, we confirmed differential regulation of key regulators in 426 defence-associated phytohormone pathways.

428 ROS production in S. chilense is highly polymorphic

429 ROS is one of the important key regulators in basal immune responses, and regulators of the 430 ROS pathway were differentially expressed in the RNAseq data (Table S5). Thus, we tested 431 ROS production in 83 genetically distinct *S. chilense* plants upon elicitation with laminarin. 432 Maximum ROS production upon laminarin elicitation was significantly different between the 433 populations (Figure 2A, Table S6). The highest average ROS maximum was recorded in 434 LA3111 and the lowest average in LA3786.

435 When grouping the populations in geographical region and looking at the overall average in 436 ROS maxima the southern highlands group had the highest ROS production and the northern 437 group had the lowest (Figure S2). We also found significant differences in ROS maxima in 438 within the individual populations for 8 out of 9 populations (Table S7), with some plants 439 showing high ROS production and others showing no detectable ROS production upon 440 elicitation with laminarin (Figure 2B and Figure S3).



441

442 Figure 2: ROS accumulation in the leaf discs from Solanum chilense measured from 0-180 minutes 443 upon elicitation with laminarin (1mg/ml). A) Overview for each of the populations. Each boxplot 444 represents a populations, each black dit the mean measured value for one plant, obtained from three or 445 four individual repetitions, as explained in B, All the significance data is highlighted in supplemental 446 material. B) examples highlighting the within-population diversity for for three of nine populations. 447 Each box plot represents an individual plant per population with data from one leaf disc represented 448 as one data point accounting up to ten to twelve leaf discs per plant. Individual measurements were 449 performed on three different dates (n=3-4 each date; $3x_3(4)=10(12)$ leaf discs per plant), each median 450 of the boxplot B represent single datapoints of A in corresponding population plant datapoints. C. Differences in ROS kinetics for the three highlighted populations from panel B. X-axis are minutes 451 452 post treatment. Y-axis shows relative luminescence unit (RLU). Colors of the boxplots or header bars 453 represent the geographical location of the population. Extended panels like B and C for all 454 populations can be found in Figure S3 and S4. Each data poiunt is a median similar to panel B and 455 the error bar represent standard error to have a clear visulatization of the different plants.

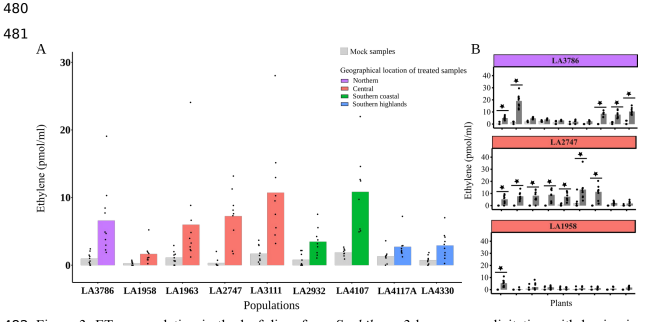
456 We also observed variation in the kinetics of the ROS production, with plants in some 457 populations not showing a clear single peak, but rather a longer-lasting ROS production. This 458 phenomenon appeared more common in the southern populations and occurs most frequently 459 in southern populations (Figure 2C and S4).

460 To confirm the specificity of the observed ROS production towards laminarin, and to show 461 that lack of observed ROS burst does not result from a general ROS signalling impairment in 462 the plants, we further tested ROS production after elicitation with the bacterial PAMP peptide 463 flg22 in all plants from population LA4330 (Figure S5). This revealed variable ROS 464 production upon elicitation with flg22. Moreover, there appears to be no apparent correlation 465 between the strengths of flg22- and laminarin-triggered responses. Some plants showed no 466 flg22 response and a clear laminarin response, or vice versa and some plants showed 467 responses in similar intensity. This suggests that the observed differences in some plants are 468 elicitor-specific variation and not a general effect of ROS production ability or defence 469 signalling pathways.

470

471 ET accumulation upon laminarin treatment is low in southern highland populations

472 To evaluate the role of phytohormones in the resistance differences observed in *S. chilense*, 473 we first looked into ET production. Upon elicitation with laminarin, we observed that plants 474 showed significant differences in ET production as compared to mock-treated samples 475 (Figure 3, Table S8 and S9). Out of 83 plants tested, we found significantly induced ET 476 production upon elicitation in 39 plants. Plant 02 from LA1963 showed a clear ET response 477 and so do several plants from population LA3111, confirming our RNAseq and gene 478 expression qPCR results above (Figure S1 and Table S9). Hence, differential ET-pathway 479 gene expression in these plants leads to laminarin-elicited ET accumulation.



482 Figure 3: ET accumulation in the leaf discs from *S. chilense* 3 hours upon elicitation with laminarin 483 (1mg/ml) and mock (milliQ H₂0). A) Each bar pair, (light grey and colored) represents an population. 484 Each bar schows the mean of the population each dot represents the mean of one plant from three

485 individual repetitions (as in B), All the significance data is highlighted in supplemental material. B) 486 Each bar pair (light grey and dark grey) represents an individual from the population. Each bar shows 487 mean of 7-9 data points which represent 7-9 samples measurements performed on three different dates 488 (n=2-3 samples each date), each sample contained three leaf discs. Significantly different ET 489 accumulation in laminarin treated samples from the mock treated samples in an individual is 490 represented with the star on the bar pair (ANOVA with post hoc Tukey tests on complete dataset). Y-491 axis shows ET accumulation in pmol/ml headspace of the samples. Each panel in B shows different 492 population and colors represent the geographical location of the population. Panels for all additional 493 populations can be found in Figure S6A.

494

495 We find significant differences in ET accumulation between the populations (Figure 3A, 496 Table S10). Looking at the populations based on their geographical locations shows that 497 overall average of ET accumulation was lowest in the southern highlands group (Figure 498 S2B). Within populations, we observed that the number of plants that significantly differ in 499 ET response varied dependent on the population. Population LA3786 showed the most 500 differences between individual plants and LA2747 was the most uniform, population LA1958 501 showed nearly no ET response (Figure 3B, Table S11).

502

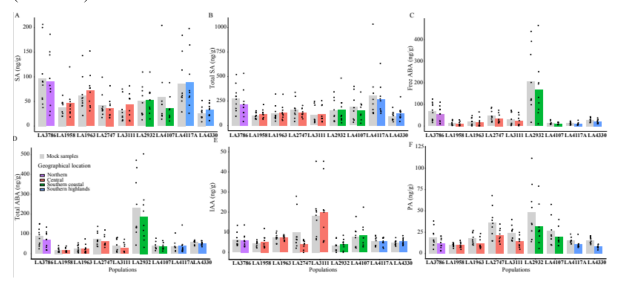
503 Populations show high diversity in the basal level of phytohormones

504 Next, we looked into the production of two important defence-related phytohormones and 505 their derivatives, for which we detected expression of key regulators in our RNAseq data, JA 506 and SA, as well as other phytohormones and their derivatives that are known to be involved 507 in stress responses: ABA, PA, DPA and IAA.

508 With our method, we were unable to detect quantifiable amounts of JA in any of the samples 509 tested, but were able to quantify both free and total SA in the basal state and after elicitation. 510 We observed that most populations did not show strong differences in the levels of SA (free 511 and total) after elicitation with laminarin (Figure 4A and B ,Figure S6 B and C and Table S8). 512 Four plants form an exception: two showed a higher amount of free SA (LA3111 plant 05 513 and LA4330 plant 05), and two showed a lower amount of free SA (LA2932 plant 12 and 514 LA4107 plant 12) when compared to the mock-treated samples (Figure S6B). These 515 differences did not correspond with those plants being more resistant or susceptible than 516 other plants in the corresponding populations.

517 Interestingly, we measured significant differences in basal levels of free and total SA within 518 and between populations. The correlation coefficient between free and total SA content is 519 0.46 (*p*-value=1.10E-05) (Table S12), therefore we treated free and total SA independently in 520 further analyses. Both basal SA levels (free and total) were significantly different between the

521 populations (Table S13). As with ROS and ET responses, we also found significant 522 differences within the populations for basal levels of both free SA and total SA content 523 (Table S14).



525 Figure 4: Phytohormone measures in the leaf discs from *S. chilense* 3 hours upon elicitation with 526 laminarin (1mg/ml) and mock (milliQ H_20). Each bar pair (light grey and colored) represents a 527 population. Each dot represents the mean of a single individual measured with at least three 528 independent repetitions. Results for each individual plant can be found in Figure S6. Y-axis shows 529 phytohormone accumulation in ng/g of the samples. Colors represent the geographical location of the 530 population.

531

We also measured ABA, PA, DPA and IAA, which are described to be important for biotic stress responses and pathways of several of these hormones influence each other. We did not detect DPA in our samples. We detected basal levels for the phytohormones ABA (free ABA, Figure 4C, Figure S6D and total ABA, Figure 4D, Figure S6E), IAA (Figure 4E, Figure S6F) and PA (Figure 4F, Figure S6G),. Laminarin treatment did not significantly change the level of these phytohormones except for PA (Table S8). For PA we observed a significantly lower amount after treatments when compared to the basal level in plants (Figure 4F, Figure S6G and Table S15). Although, in general basal levels of PA and PA levels upon elicitation were highly correlated (Pearson's correlation coefficient of 0.74, *p*-value=2.2E-016) (Table S12). We observed a higher amount of basal levels of free and total ABA in LA2932 as compared to other populations, whereas IAA was higher in LA3111. The levels of all these phytohormones show significant differences between and within populations (Table S16 and table S17, respectively) in an independent manner (Table S12). Looking at the data of all the tested populations based on geographic location, we found higher levels of basal PA in the

546 southern coast and IAA was higher in the central region whereas SA (free and total) was high547 in the north and ABA levels were higher in northern and south coast populations (Figure S2)..548

549 Multiple defence responses correlate with observed resistance phenotypes

To assess whether the individual defence responses measured in the plants can be associated with *S. chilense* resistance properties, we looked for correlations with previously generated data on the frequency with which *P. infestans* can infect *S. chilense* leaflets, the so-called infection frequency (IF) (Kahlon *et al.*, 2021). We found no correlation between the observed ROS maxima and the IF observed with *P. infestans* (Pearson's correlation coefficient of 0.09, *p*-value=0.37, Table 1), whereas we found a significant negative Pearson's correlation of IF with ET accumulation (-0.36, *p*-value=0.0008; Table 1). The Pearson's correlation of IF with basal levels of PA also showed a negative correlation (-0.2443413, *p*-value=0.026; Table 1), whereas we observed no strong or significant (*p*-value<0.05) correlation for SA , ABA and IAA (Table 1).

560

562

563 Table 1: Pearson's correlation of measured potential immunity-related factors at basal levels and upon 564 elicitation with laminarin (1mg/ml) with the infection frequency (IF) of same plants upon inoculation 565 with *P. infestans* Pi100 published in Kahlon *et al.* (2021). The correlation is shown for the all the 566 measured components. Significant correlation is highlighted in green.

567

Parameter compared	Pearson's correlation coefficient	<i>p</i> -value
IF~ROS	0.10	0.3748
IF~ET	-0.36	0.0008
IF~Free SA (Basal)	-0.05	0.6290
IF~Total SA (Basal)	-0.15	0.0690
IF~Free IAA (Basal)	-0.13	0.2727
IF~Free ABA (Basal)	-0.12	0.2737
IF~Total ABA (Basal)	-0.12	0.2419
IF~PA (Basal)	-0.24	0.0260

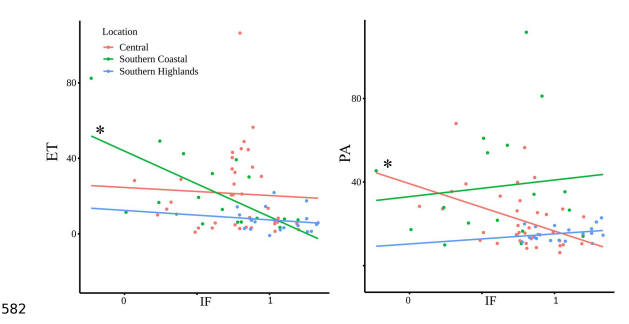
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571 The dominance of individual defence responses differs geographically

572 All measured components showed geographical trends at basal and induced levels (Figure S2, 573 as well as S3, S4, S6). This supports that the plants' genotypes rather than the common 574 experimental environment was the driver of metabolic differences between the plants. We 575 hypothesize that different populations have adapted different defence strategies due to 576 adaptation to specific climatic niches. Thus, to confirm the possible larger effect any of 577 measured components in certain geographical regions, we calculated the correlation 578 coefficient for ET and PA for each of the geographical groups of *S. chilense*. We found that 579 the effect of ET is most strongly correlated with resistance in the coastal populations, 580 whereas PA showed the strongest correlation to resistance in the central group (Figure 5 and 581 Table S18).

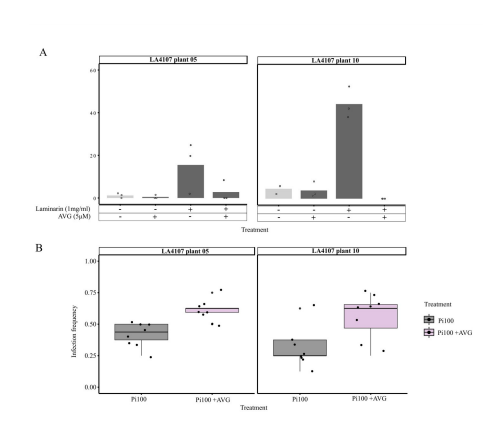


583 Figure 5: Correlations between the *P. infestans* infection frequency (IF, X-axis) and the measured ET 584 or PA response phytohormone accumulation in ng/g of the samples or pol/ml respectively (Y-axis). 585 Each dot represents the mean value of an individual plant from three individual repetitions. Infection 586 frequencies were obtained from and independent experiment from the same plants as presented in 587 Kahlon et al (2020). IF at zero indicates plants that are fully resistant and IF at one shows 100% 588 infection rate upon inoculation. The pearson correlations were calculated per geographic group. The * 589 indicates the significant correlations for the central and southern coastal populations for PA and ET 590 respectively.

591

592 ET plays a role in defence response in southern coast tested individuals

To confirm the contribution of ET to resistance in the coastal populations, we selected two plants from southern coastal population LA4107, one with relatively high ET production and another with relatively moderate ET production, with low and medium-high scores from the infection frequency spectrum, respectively. To verify the role of ET on the resistance outcome, we used AVG, a well-established chemical inhibitor, to halt the ET production in the selected plants and tested the ET accumulation after laminarin treatment. AVG was successfully able to inhibit the ET production up to 100% in the plant samples (Figure 6A). After inoculation with *P. infestans* isolate Pi100, the plants indeed showed higher succeptibility when they were pre-treated with AVG as compared to control plants, confirming the positive role of ET in basal resistance in this population (Figure 6B).



603

604

605 Figure 6: ET inhibition assay on LA4107 plant 05 and plant 10: ET accumulation in the the leaf discs 606 from *S. chilense* LA4107 plant 05 and 10, 3 hours upon elicitation with laminarin (1mg/ml), AVG 607 (5μ M), laminarin (1mg/ml) + AVG (5μ M), and mock (milliQ H₂0). Light and dark grey crossbar pair 608 represents plant with and without treatment with laminarin (1mg/ml). Each crossbar is the mean of 609 three samples measurement. Y-axis shows ET accumulation in pmol/ml air of the samples. b) 610 Detached leaf infection assay of LA4107 plant 05 and 10 upon drop inoculation with *Phytopathora* 611 *infestans* Pi100 (3000 sporangia/ml) with and without AVG (5μ M) treatment. Y-axis represents 612 infection frequency which is the ratio of infected leaflets divided by inoculated leaflets. Each dot 613 represents the ratio from one leaf, the red dot represents the mean value and the *p*-value is shown on 614 the top of the boxplot.

615

616 **Discussion**

617 We previously used natural populations of *S. chilense* to show intraspecific variation in 618 resistance against *P. infestans* (Kahlon *et al.*, 2021). In this study, we evaluated several key 619 components of basal defence responses in the same plants to explore molecular cues behind 620 the previously observed phenotypic variation.

621 In order to reliably and reproducibly study defence components in this polymorphic plant 622 species, and to rule out variation arising during the preparation of pathogen biological 623 material, we used the glucan elicitor laminarin. Laminarin has been previously reported to 624 activate basal immune responses such as ROS production, calcium influx and MAPK 625 activation in members of the Solanaceae family (Meénard *et al.*, 2004; Wanke *et al.*, 2020).

626 We observed significant overlaps in DEGs in *S. chilense* central population LA3111 upon 627 elicitation with laminarin and infection with *P. infestans*. The majority of these genes are 628 known for involvement in defence responses. We further showed differences in transcript 629 levels via qPCR of key regulators of defence-related phytohormones after laminarin 630 treatment in a plant of a different central population, LA1963. This suggests that laminarin 631 can be used as a proxy for evaluating early basal immune responses activation in *S. chilense*.

The RNAseq data of both *P. infestans* and laminarin treatments revealed regulation of homologs of several previously identified genes known in major defence pathways, like ROS broduction and both SA and JA signalling. These basal immunity components have been in the been based in *P. infestans* resistance in different Solanaceous plant species. In cultivated tomato, reduced accumulation of ROS results in enhanced resistance against *P. infestans* (Cui *et al.*, 2016). Higher SA levels have a positive effect on *P. infestans* resistance in *S. tuberosum* (Halim *et al.*, 2007). Both, SA and ET contribute to resistance in *N.* benthamiana (Shibata *et al.*, 2010), and higher levels of JA and interplay with SA were do observed in resistant cultivars of *Capsicum annuum* (Ueeda *et al.*, 2005). We observed high intraspecific diversity in the above-mentioned components early after elicitation with laminarin, and at basal levels. Large-scale intraspecific variation in basal immunity has also been reported on a transcriptional level in *A. thaliana* accessions, upon elicitation with the bacterial PAMP flg22 (Winkelmüller *et al.*, 2021).

645 Surprisingly, we did not find a strong correlation between the amount of ROS produced in a 646 plant after elicitation with laminarin and its resistance properties. ROS production upon biotic 647 stress has often been considered as a hallmark of successful recognition of pathogens and the 648 activation of defence (Torres, 2010). ROS production linked to the perception of flg22 is 649 often taken as an indicator for resistance against bacterial *Pseudomonas* spp. pathogens (e.g. 650 in *A. thaliana*; Smith and Hesse, 2014 and in tomato; Roberts *et al.*, 2019). Our study shows 651 that laminarin-triggered ROS production in *S. chilense* cannot be used to estimate the basal 652 resistance against *P. infestans*. Similarly, we also found no correlation between laminarin-653 triggered SA production or basal SA levels and *P. infestans* resistance. Thus, these individual 654 defence components triggered by laminarin either have a rather limited contribution to the 655 observed *P. infestans* resistance in the populations, or ROS and SA are not directly involved 656 in *P. infestans* resistance in *S. chilense*. On the other hand, laminarin has been shown to 657 induce the ET pathway, but only sulphated laminarin (β-1,3 glucan sulfate) can induce the 658 salicylic acid signaling pathway in *N. tabacum* and *A. thaliana* (Meénard et al. 2004). In our 659 RNAseq analysis, we do see more DEGs when plants are treated with *P. infestans* as

660 compared to laminarin. In the future, it would be interesting to evaluate the effects of 661 sulphated laminarin and other known PAMPs from *P. infestans* in order to dissect the defence 662 responses further.

663 Our data does support that resistance observed in our plant species can be correlated to 664 different components in the plants: induced ET and also basal levels of the phytohormone 665 PA. This is in line with the hypothesis that basal defence is regulated by a complex network 666 of interacting components from plants and pathogens (Windram and Denbi 2015, Kahlon 667 and Stam, 2021a). We also observed that the strength of the correlation is dependent on the 668 geographical region from which the plants originated. Calculations per population would be 669 even more interesting, but due to the limited number of plants per population, these 670 calculations would lack statistical power.

671 It can be assumed that in each populations multiple components play important roles, but that 672 due to sample size limitations, these effects were not picked up. The generation of 673 generalized linear mixed models, testing the combined effect of multiple components would 674 be desirable in this context, though this would require a lot of additional data.

675 It has previously been shown that ROS production leads to SA production in a feed-forward 676 loop in defence responses in *Arabidopsis* (reviewed by Herrera-Vásquez *et al.*, 2015). We did 677 not observe such correlation among ROS production and SA production at early time points, 678 nor did we observe a correlation between SA and ET production as observed in tomato 679 resistance against the fungal pathogen *Fusarium oxysporum* (Di *et al.*, 2017) . Interestingly, 680 the suppression of ABA biosynthesis and activation of ET biosynthesis upon copper ions 681 treatment enhances resistance against *P. infestans* in potato seedlings (Liu *et al.*, 2020b), 682 whereas in our system ET positively contributed to resistance observed against *P. infestans*.

In our assays, ET had the strongest role in early defence. ET has previously been described in association with various defence responses. In *A. thaliana*, Resistance to Powdery Mildew 8 (RPW8)-mediated defence response is regulated by a ET-mediated feedback loop (Zhao *et al.*, 2021). For Solaneceous species, the activation of defence-related genes in *P. infestans* resistant potato cultivars upon exogenous ET treatment has also been reported in a recent transcriptome study, by Yang *et al.*, (2020) and laminarin triggers the expression of ETdependent defence genes in *N. tabacum* (Meénard *et al.*, 2004). A positive role of ET production has been reported in relation to resistance to *C. fulvum* in tomato plants carrying corresponding resistance genes against a specific *C. fulvum* race (Hammond-Kosack *et al.*, 1996). Another study showed that ET is also involved in resistance to the fungal pathogen *B*.

693 *cinerea* and certain wound responses in tomatoes, with no clear role of JA or SA observed 694 (Díaz *et al.*, 2002).

695 We further showed geographical variation in basal and induced levels of each component and 696 expect that the role of each component might differ between populations due to variation in 697 habitats. Interestingly, the role of ET was stronger in the coastal populations and 698 experimentally verified with ET inhibition assays in plants from coastal population LA4107. 699 The stronger association of ET and resistance specifically in the southern coastal populations 700 could be a result of specific adaptation processes in these populations. This could potentially 701 reflect an added benefit of the development of stronger ET signalling in these populations as 702 a result of specific habitat adaptation e.g. to deal with potential abiotic stresses, like salt or 703 temperature. General temperature dependency of defence regulation and the involvement of 704 phytohormone signalling has been shown for both cold (Wang et al., 2019) and heat stress 705 (Huang et al., 2021). ET has been shown to be a crucial phytohormone when it comes to 706 coping with salinity stress in plants (Riyazuddin et al., 2020). The positive effects of ET in 707 salt tolerance have been illustrated in A. thaliana (Yang et al., 2013) and Zea mays (Freitas et 708 al., 2018). In a study by Kashyap et al., (2020), S. chilense plants under salt stress coped 709 better than cultivated S. lvcopersicum due to a better anti-oxidant system. We also observed 710 high basal levels of the phytohormone ABA in the coastal population LA2932 (Figure 4C-D). 711 ABA is highlighted to be an important phytohormone for abiotic stress tolerance including 712 salinity stress (reviewed by Zhu, 2002 and Ng et al., 2014).

The genotype-to-phenotype linkages in systems biology are complex. In a diverse panel of wild and domesticated tomatoes basal resistance against the generalist fungal pathogen *Botrytis cinerea* has been reported to be dependent on the interaction of multiple loci among both host and pathogen (Soltis *et al.*, 2019). Here we presented a decomplexification approach, where more components can be added to understand both the triggers (by testing different elicitors) and the outcomes (by measuring more responses). As highlighted by Marshall-Colón and Kliebenstein, (2019), such future studies should be performed using large-scale metabolomics and transcriptomics analyses, to determine the key regulators underlying the measured responses and to be able to appreciate the intrinsic value of the complexity of signalling networks.

723 In our previous studies (Stam *et al.*, 2017; Kahlon *et al.*, 2021) we showed no strong signs of 724 host adaptation towards resistance to *P. infestans*, as resistance shows no clear geographical 725 pattern of adaptation. However, our current results indicate that different coping mechanisms 726 are present in each of these populations, possibly due to specific adaptation to the niches that

727 the plants inhabit in each region. Examples of such specific adaptations have also been 728 observed for other host-parasite interactions. Populations of *Eruca vesicaria* (syns. *Eruca* 729 *sativa*, wild rocket) from Mediterranean and desert habitats showed activation in defence 730 responses via two different mechanisms when challenged with generalist herbivore 731 *Spodoptera littoralis*. Mediterranean plants showed accumulation of glucosinolates and desert 732 plants showed induced levels of a specific protease inhibitor (Ogran *et al.*, 2016). Beevan *et 733 al.* (1993) showed immense differences in phenotypes of two populations of *Senecio vulgaris* 734 (groundsel) against *Erysiphe fischeri* and proposed different populations have evolved 735 different survival strategies against the same pathogen. In *A thaliana*, combined effects of 736 genetic variation and differences in environmental factors also shape defence-associated 737 metabolite contents (Katz *et al.* 2021)

Together our data support high complexity of *S. chilense*'s defence response to the general glucan elicitor laminarin. These responses might contribute to *P. infestans* resistance by additive or network functions. At single geographic location, certain plant hormones play a to the local abiotic environment and hormones may be key to this adaptation. The corresponding defence machinery might simultaneously undergo co-adaptation to cope with biotic stress. We speculate that plants' high connectivity between abiotic and biotic signaling results in the necessity to habitat-specifically recruit different defence pathways and that the nature of the involved hormones but also strongly by the abiotic environment and highlights the need for further population-scale studies on pathogen resistance mechanisms (Kahlon and Stam 2021).

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750

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763

764 Author contributions

765 Conceptualization: RS, PSK, CD, RHü and JB; Investigation: PSK, AF, MM, MO, MG and766 RHa; Data interpretation and evaluation: PSK, MM, AF, MO, RHü and RS; Writing and Data767 representation: PSK and RS. All authors reviewed and approved the manuscript.

768

769 Conflicts of interests

770 The authors declare that no competing interests exist.

771

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776 Data Statement

777 Analytical results are included in the supplementary data files. Raw sequence data (Illumina 778 reads) are uploaded to NCBI SRA and available under PRJNA746795. All scripts used for 779 the analyses, as well as all intermediate data files (e.g. FeatureCount output files) can be 780 found on Zenodo (10.5281/zenodo.5101308)

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