# The wild tomato species *Solanum chilense* shows local variation in pathogen resistance between geographically distinct populations (#13374)

First submission

Please read the **Important notes** below, and the **Review guidance** on the next page. When ready **submit online**. The manuscript starts on page 3.

Important notes						
<b>Editor</b> Erica Goss						
Files	6 Figure file(s) 1 Table file(s) 1 Raw data file(s) 1 Other file(s) Please visit the overview page to <u>download and review</u> the files not included in this review pdf.					
Declarations	No notable declarations are present					

### Review guidelines

Please in full read before you begin



#### How to review

When ready **submit your review online**. The review form is divided into 5 sections. Please consider these when composing your review:

- **1. BASIC REPORTING**
- 2. EXPERIMENTAL DESIGN
- **3. VALIDITY OF THE FINDINGS**
- 4. General comments
- 5. Confidential notes to the editor
- You can also annotate this **pdf** and upload it as part of your review

To finish, enter your editorial recommendation (accept, revise or reject) and submit.

#### **BASIC REPORTING**

Clear, unambiguous, professional English language used throughout.
 Intro & background to show context. Literature well referenced & relevant.
 Structure conforms to Peerl standard, discipline norm, or improved for clarity.
 Figures are relevant, high quality, well labelled & described.
 Raw data supplied (See Peerl policy).

#### VALIDITY OF THE FINDINGS

- Impact and novelty not assessed. Negative/inconclusive results accepted. *Meaningful* replication encouraged where rationale & benefit to literature is clearly stated.
  - Data is robust, statistically sound, & controlled.

#### **EXPERIMENTAL DESIGN**

 sh
 Original primary research within Scope of the journal.

 Image: Short Scope of the journal.
 Research question well defined, relevant & meaningful. It is stated how research fills an identified knowledge gap.

 Image: Short Scope of the journal.
 Rigorous investigation performed to a high technical & ethical standard.

 Image: Short Scope of the journal.
 Rigorous investigation performed to a high technical & ethical standard.

 Image: Short Scope of the journal research question to replicate.
 Methods described with sufficient detail & information to replicate.

 Image: Short Scope of the journal research question & limited to supporting results.
 Speculation is welcome, but should be identified as such.

The above is the editorial criteria summary. To view in full visit <u>https://peerj.com/about/editorial-</u> criteria/

# The wild tomato species *Solanum chilense* shows local variation in pathogen resistance between geographically distinct populations

Remco Stam Corresp., 1 , Daniela Scheikl 1 , Aurélien Tellier 1

<sup>1</sup> Section of Population Genetics, Technical University of Munich, Freising, Germany

Corresponding Author: Remco Stam Email address: stam@wzw.tum.de

Wild tomatoes are a valuable source of disease resistance germplasm for tomato (Solanum *lycopersicum*) breeders. Many species are known to possess a certain degree of resistance against certain pathogens, however evolution of resistance traits is yet poorly understood. For some species, like Solanum chilense, both differences in habitat and within species genetic diversity is very large. Here we aim to investigate the occurrence of spatially heterogeneous coevolutionary pressures between populations of S. chilense. We investigate the phenotypic differences in disease resistance within S. chilense against three common tomato pathogens (Alternaria solani, Phytophthora infestans and a *Fusarium sp.*) and confirm high degrees of variability in resistance properties between selected populations. Using generalised linear mixed models, we show that disease resistance does not follow the known demographic patterns of the species. Models with up to five available climatic and geographic variables are required to best describe resistance differences, confirming the complexity of factors involved in local resistance variation. We confirm that within S. chilense, resistance properties against various pathogens show a mosaic pattern and do not follow environmental patterns, indicating the strength of local pathogen pressures. Our study can form the basis for further investigations of the genetic traits involved.



1

- The wild tomato species Solanum chilense shows local variation in pathogen resistance
- 2 3 between geographically distinct populations
- 4
- 5 Remco Stam\*, Daniela Scheikl, Aurélien Tellier
- 6 Section of Population Genetics, Technische Universität München, Liesel-Beckmann Strasse 2,
- 7 85354 Freising, Germany
- 8 \*Author for Correspondence: Remco Stam: stam@wzw.tum.de

9

10

11

12

#### 14 Abstract

15

16 Wild tomatoes are a valuable source of disease resistance germplasm for tomato (Solanum 17 lycopersicum) breeders. Many species are known to possess a certain degree of resistance 18 against certain pathogens, however evolution of resistance traits is yet poorly understood. For 19 some species, like Solanum chilense, both differences in habitat and within species genetic 20 diversity is very large. Here we aim to investigate the occurrence of spatially heterogeneous 21 coevolutionary pressures between populations of S. chilense. We investigate the phenotypic 22 differences in disease resistance within S. chilense against three common tomato pathogens 23 (Alternaria solani, Phytophthora infestans and a Fusarium sp.) and confirm high degrees of 24 variability in resistance properties between selected populations. Using generalised linear mixed 25 models, we show that disease resistance does not follow the known demographic patterns of 26 the species. Models with up to five available climatic and geographic variables are required to 27 best describe resistance differences, confirming the complexity of factors involved in local 28 resistance variation. We confirm that within S. chilense, resistance properties against various 29 pathogens show a mosaic pattern and do not follow environmental patterns, indicating the 30 strength of local pathogen pressures. Our study can form the basis for further investigations of 31 the genetic traits involved.

32

33

#### 34 Keywords

Host pathogen interaction, resistance, wild tomatoes, alternaria, fusarium, phytophthora, localvariation

37

#### 39 Background

40

41 In nature, plants are exposed to a wide range of pathogens and pests. While in most cases the 42 plants appear non-specifically resistant against these threats, drastic or recurrent epidemics do 43 occur (Thrall et al. 2001a, Soubeyrand et al. 2009) and variability in specific resistance to 44 pathogens is observed (Thrall et al. 2001b, Salvaudon et al. 2008). Understanding how 45 reciprocal co-adaptation of hosts and pathogens maintains such diversity has been a key 46 guestion in theoretical and empirical evolutionary biology. Theoretically, negative direct 47 frequency-dependent selection (ndFDS) is shown to be a necessary condition to maintain long-48 term stable diversity for resistance in plants and infectivity in pathogens (Tellier and Brown 49 2007). Seed banking, perenniality or polycyclic disease can generate ndFDS, while costs of 50 resistance and infectivity (virulence) are necessary but not sufficient for stable long term 51 polymorphism to occur (Tellier and Brown 2009, Brown and Tellier 2011). Another factor often 52 suggested to maintain diversity is the spatial structure of host and pathogen populations. Spatial 53 structure and migration of hosts and pathogens as well as population sizes and genetic drift 54 generate patterns of local adaptation over space and time (Thrall and Burdon 2002, Gandon 55 and Michalakis 2002). However, a spatial structure with homogeneous environment does not 56 generate ndFDS (Thrall et al. 2002a, Tellier and Brown 2011) Stable long term polymorphism is 57 favoured by spatially heterogeneous environments across which the prevalence and severity of 58 disease or the costs of resistance and infectivity may differ (Gavrilets and Michalakis 2008, 59 Moreno-Gamez et al. 2013). 60 From an ecological perspective, and based on the classic disease triangle from plant pathology 61 (Agrios 2005) the outcome of species interactions are mediated by the abiotic and biotic 62 environment. The influence of the environment generates therefore spatial and temporal

- 63 variation in evolutionary and coevolutionary dynamics (Thompson 2005), and increasing
- 64 evidence for geographical variation in coevolutionary dynamics and patterns of local adaptation

are found in microcosm experiments (Forde et al. 2004, Vogwill et al. 2009, Lopez-Pascua et al.2010)

67 Nevertheless, few field systems exist to study and document the coevolution of plants and their 68 pathogens occurring at short time scales and across several populations. One example is the 69 wild flax – flax rust pathosystem, where local adaptations have been observed and the most 70 resistant varieties of flax generally harboured more virulent strains of rust (Thrall et al. 2002a, 71 Thrall and Burdon 2003). Similarly, the local adaptation of powdery mildew Podosphaera 72 plantaginis to Plantago lanceolata populations spread over different islands off the coast of 73 Sweden showed virulent strains to co-occur with more resistant plants (Laine 2005, Soubeyrand 74 et al. 2009). In the latter plant-pathogen system, several mechanisms theoretically proposed to 75 generate ndFDS have been shown to originate from the environmental heterogeneity across 76 populations: 1) GxGxE interactions (host genotype x pathogen genotype x environment, for example (Laine 2 2) heterogeneity in disease incidence and prevalence determining thus 77 78 epidemiological pressures (Soubeyrand et al. 2009) and co-infection (Susi et al. 2015), and 3) 79 different strength of connectivity between populations accelerating or decelerating the speed of 80 coevolution across the landscape (Jousimo et al. 2014). These factors are thus expected to 81 promote and facilitate long term polymorphism at resistance and infectivity loci without 82 unrealistic costs of these alleles. Here we aim to investigate the occurrence of spatially 83 heterogeneous coevolutionary pressures between populations of Solanum chilense, a 84 solanaceous wild species, and several pathogens in a relatively small geographical space which 85 exhibits large variation in habitat quality and abiotic environmental factors.

86

Wild Solanum species are in general particularly good model species to study between and
within species variation, because they occur in diverse geographic and climatic habitats and
have a very well studied demography and known evolutionary history (Städler et al. 2005, 2008,

Tellier et al. 2011) Additionally, several studies exist that suggest that at least bacterial resistance-associated genes are under selective pressure (Rose et al. 2005, 2007, 2011). *S. chilense* is native in South America, ranging from southern Peru to central Chile, in a broad range of habitats. *S. chilense* populations have been found from coastal regions, even in slightly alkaline environments, all the way to high altitude (>3000 m) mountain regions. It has been found in extreme dry habitats on the border of the Atacama dessert, as well as near rivers and creeks (Peralta et al. 2008).

97 S. chilense most likely originated with its sister species S. peruvianum, somewhere in south 98 Peru and then migrated south (Städler et al. 2008). A study of the species' demography found 99 four genetically distinct subgroups; one in the north of the range, one in the central region and 100 two in the south (one on the coast and one at high altitudes). Interestingly, the two southern 101 groups are, even though geographically close to each other, more related to the central group 102 than to each other, possibly due to the separating effect of the extremely arid Atacama desert 103 (Böndel et al. 2015), In addition, S. chilense shows clear climatic adaptations. Populations from 104 drier regions are responding faster to drought (Fischer et al. 2013) and individual populations 105 found at high altitudes (>3000 m) show higher freezing tolerance (Nosenko et al. 2016) S 106 chilense has also been the source of resistance loci against the fungus Verticilium dahliae 107 (Tabaeizadeh et al. 1999) and against various viruses (Griffiths and Scott 2001, Ji et al. 2007, 108 Verlaan et al. 2013). Seeing that S. chilense occurs in such a wide range of habitats and that 109 the species shows specific signs of local climatic adaptations, we wondered whether we could 110 find variation for pathogen resistance as well.

111

Since no exace the exist about the co-occurrence of wild pathogens and *S. chilense*, we chose to test *S. chilense* disease resistance properties with three widely studies and economically relevant pathogens, *Alternaria solani*, Phytophthora *infestans* and a *Fusarium* sp.

A. solani causes early blight and is amongst the most destructive diseases of tomato in tropical and subtropical regions, leading to yield losses of up to 80% in certain regions. *A. solani* has been found in central Peru and is known to cause disease not only on potato - its main host but also on many other nightshades, including tomato (Song et al. 2011, Kumar et al. 2013). In addition, previous work has shown that *A. solani* resistance can be studied using detached leaf assays (Chaerani and Voorrips 2006, Chaerani et al. 2007).

121 Fusarium spp are pathogens that cause very severe disease symptoms on a very wide range of 122 host plants that span almost the entire globe (Agrios 2005) Two Fusarium spp are on the top10 123 most important fungi in plant pathology (Dean et al. 2012) The F. oxysporium species complex 124 comprises over 100 formae specialis that all infect specific hosts, including tomato (Michielse 125 and Rep 2009). It is widely used to study molecular and genetic mechanisms involved in plant 126 pathogen interactions (Houterman et al. 2008, Ma et al. 2013) and even though it is generally 127 reported to be a vascular pathogen, it has regularly been successfully deployed in detached leaf 128 infection assays (e.g. (Kavroulakis et al. 2007)).

129 *Phytophthora infestans* is an oomycete that causes late blight on potato and tomato. In potato 130 alone the damage amounts up to \$1 bn annually (Haverkort et al. 2009)Due to its economic 131 value and the vast amount of molecular and genetic research performed on it, it is considered 132 the most important oomycete plant pathogen (Kamoun et al. 2015). Like the other two 133 pathogens used in this study, P. infestans strains have been sampled in parts of the natural 134 habitat of S. chilense (Perez et al. 2001). The strain EC1 that we used has its origin in Ecuador 135 and is particularly relevant for agriculture as it is a rather aggressive strain that is capable of 136 overcoming certain novel genetic resistances (Foster et al. 2009, Nowicki et al. 2011).

137

Here we test the resistance of different *S. chilense* populations from three different regions in
Chile and Peru, one central region and two southern regions, one coastal and one mountainous

140 (see Fig1b) against the above mentioned pathogens. These group resemble very distinctive

- 141 habitats and can thus be used to investigate whether we see differences in infection rate
- 142 throughout the range of the species. We also test whether these differences show a linear
- 143 pattern when tested against geographical and climatic variables (e.g. north more resistant, high
- 144 precipitation more resistant) or whether a multitude of factors leads to specific local adaptations
- 145 to each of the three pathogens.

147 Methods 148 Plant growth 149 Seed batches were obtained from the tomato genomics resource centre (TGRC, Davis, USA). 150 We grew seven different Solanum chilense populations (accession numbers LA1963, LA2931, 151 LA2932, LA3111, LA4107, LA4117 and LA4330) consisting of 10 different plants each and one 152 Solanum pennellii (LA0716) population in our glasshouse from randomly chosen seeds. The 153 plants were grown with 16h light and a minimum temperature of 18°C. Mature plants were cut 154 back at a biweekly interval to assure young leaves of similar age were available at all times for 155 all populations 156 157 Pathogen propagation and spore production 158 Alternaria solani 159 A. solani strains B055 and St108 were obtained from the chair of Phytopathology at the TUM (Munich, Germany) and cultivated on SN plates (at 22°C, 12h UV-A light, 12h darkness 160 161 (induction of sporulation) and 85% humidity for 3 weeks. We harvested the spores with ddH<sub>2</sub>0 162 by scratching the mycelium with off the agar. The solution was filtered through 4 layers of mesh 163 and diluted to a concentration of 5000 spores per ml. Each leaflet was infected with a 10µl 164 droplet. 165 Phytophthora infestans 166 We obtained late blight pathogen *P. infestans* strain EC1 from the James Hutton Institute 167 (Dundee, UK). It was cultivated on RyeB agar, incubated 6 days at RT in darkness, 3 days at 168 RT and daylight. We scratched the mycelium with ice cold water with a pipette tip from the plate 169 and store at 4°C until further use (up to 3 hours). The solution was diluted to 2000-3000 170 sporangia per ml and the leaflets were infected with 5µl of this solution. 171 Fusarium sp. 172 Fusarium infected lesions were identified on a few detached S. chilense leaves from our

### Manuscript to be reviewed

glasshouse. These lesions were extracted and re-cultivated for several rounds on PotatoDextrose-Agar (PDA) for clean-up. Microscopic observations and sequence analysis of a cloned
Tubulin Beta gene confirmed the genus. Once clean, the *Fusarium* was grown on PDA for a
minimum of four days at RT. Spores were harvested by adding ddH<sub>2</sub>O and aspirating the liquid.
The spores were diluted to 2x10<sup>5</sup>-5x10<sup>5</sup> spores per ml and we infected the individual leaflets
with 5µl of this solution.

179 All protocols for pathogen cultivation, including ingredients for the growth media can be found in

180 more detail on https://www.protocols.io/view/Plant-Pathogen-Cultivation-fmkbk4w

181

#### 182 Infection assays

183 To minimise the effect of variation between plants within one population, we collected leaves of 184 same age randomly from 8 to 10 plants per population and shuffled them. We then drew the 185 leaves randomly from that mix to distribute them over up to 9 boxes for each infection 186 experiment. Each box contained 16 leaves (4 rows), from four different populations and each 187 box contained different combinations of populations. Box number and leaf position were marked 188 to later rule out possible effects. To eliminate the possible confounding effect of difference in 189 surface coating composition between the different populations and remove any pathogens that 190 accumulated on the plants during the growth time in the glass house, we washed them for 10 191 seconds with 70% Ethanol to sterilize the surface and remove natural wax layers before 192 washing with ddH2O. We assured the leaf surface was dry before drop inoculation. For each 193 pathogen 16-24 leaves - about 100 leaflets - were infected for each population and the 194 experiments were repeated four times, accumulating to about 450 – 500 infection events per 195 pathogen. The Alternaria infections were done on the axial side of the leaf, Phytophthora and 196 Fusarium infections were done on the abaxial side of the leaf. The leaves were incubated at RT 197 and scored after 6 to 8 days, dependent on temperature and growth conditions in the lab.

199	Scoring and Data analysis
200	All data analysis was done using R (R foundation for statistical computing). Generalised Linear
201	Mixed Models were made using the glmer option from the package Ime4. To construct GLMM
202	we used a binomial variable (y) consisting of the number of successful and unsuccessful
203	infection events per leaf. The GLMM were constructed taking the leaf position in the box (leaf)
204	and a combination of the box number and experimental date (exp:box) into account as random
205	effects. For our first model populations names were used as fixed effects. (model1 = y $\sim$
206	accession +(1 leaf)+(1 exp:box)). For the second model, we hierarchically tested different
207	climatic and geographical parameters (e.g. model2 = y $\sim$ geographic1 + climatic1 + climatic 2 +
208	(1 leaf)+(1 exp:box)). Pairwise comparisons were examined using an implementation of Tukey
209	Honest Significant Difference test as provided by function glht from the the R package
210	multcomp. The boxplots were drawn using the package ggplot2 and the heatmap using gplots.
211	All packages are available through CRAN.
212	
212 213	Distribution map and geographical characteristics
	<b>Distribution map and geographical characteristics</b> Geographical data for all populations were obtained from the Tomato Genome Resource
213	
213 214	Geographical data for all populations were obtained from the Tomato Genome Resource
213 214 215	Geographical data for all populations were obtained from the Tomato Genome Resource Centre. Climatic data were extracted from the <u>http://worldclim.org/</u> database. The species
<ul><li>213</li><li>214</li><li>215</li><li>216</li></ul>	Geographical data for all populations were obtained from the Tomato Genome Resource Centre. Climatic data were extracted from the <u>http://worldclim.org/</u> database. The species distribution map was drawn using the maps package in R. All geographic and climate data used
<ul> <li>213</li> <li>214</li> <li>215</li> <li>216</li> <li>217</li> </ul>	Geographical data for all populations were obtained from the Tomato Genome Resource Centre. Climatic data were extracted from the <u>http://worldclim.org/</u> database. The species distribution map was drawn using the maps package in R. All geographic and climate data used
<ul> <li>213</li> <li>214</li> <li>215</li> <li>216</li> <li>217</li> <li>218</li> </ul>	Geographical data for all populations were obtained from the Tomato Genome Resource Centre. Climatic data were extracted from the <u>http://worldclim.org/</u> database. The species distribution map was drawn using the maps package in R. All geographic and climate data used
<ul> <li>213</li> <li>214</li> <li>215</li> <li>216</li> <li>217</li> <li>218</li> <li>219</li> </ul>	Geographical data for all populations were obtained from the Tomato Genome Resource Centre. Climatic data were extracted from the <u>http://worldclim.org/</u> database. The species distribution map was drawn using the maps package in R. All geographic and climate data used

PeerJ reviewing PDF | (2016:09:13374:0:0:NEW 19 Sep 2016)

Manuscript to be reviewed

223 Results

224

225 S. chilense populations show different resistant properties against different pathogens 226 We selected seven populations that represent three previously described genotype groups 227 (Böndel et al. 2015). Two populations originate from the central range (LA1958, LA3111), two 228 from the coastal regions (LA2932, LA4107) and two from the southern mountainous region 229 (LA4117, LA4330). A seventh population is geographically in the middle between the southern 230 mountain and the central group (LA2931). Böndel et al. group it with the central populations, but 231 assign properties of both groups to it. Figure 1A shows the species distribution and highlights 232 the selected populations.

233

234 some wild tomato species (e.g. *S. pennellii*), thick and sticky surface coating have a dramatic
235 effect on pathogen ingress. In *S. chilense*, surface coatings are notably less thick, and resemble
236 those of cultivated tomato, however to minimise the effect of difference in coating, as well as to
237 sterilise the leaves, we washed all leaves briefly in 70% ethanol before infection. The effects of
238 *S. chilense su* seterilisation is noticeable during infection, but not as dramatic as with *S*239 *pennellii* (S Figure 1).

240 We infected individual leaflets for up to 16 leaves of each population per experiment with 241 Alternaria solani (str 108) and counted the occurrence of infected leaflets per leaf, as this 242 represents the success rate of the pathogen to establish itself and overcome genetic resistance. 243 Infection events, were scored as either negative (no infection or clear small necrotic lesions, 244 indicating a hypersensitive response) or positive (ranging from growth just outside the droplet 245 area up to full infection of the leaflet) (Fig 1B). We observed variation within each population. In 246 almost all instances at least one leaf was fully infected whereas another was completely 247 resistant. These outliers have large effect on the calculated mean fraction. To allow good 248 judgement we report the 1<sup>st</sup> and 3<sup>rd</sup> quantile, the median value as well as the mean value for

249	each population (Fig 2). The mean and median of the infected fractions range from 0.35 and
250	0.42 for LA3111 to 0.74 and 0.81 for LA4330 or 0.67 and 0.82 for LA2932.
251	To test the robustness of our method, we did an additional infection with a second strain of
252	Alternaria (B055). The overall infection rates are lower in this set of experiments (median of 0.54
253	compared to 0.62), however Figure S2 shows that just like for strain st108, LA3111 is the least
254	infected population with a mean of 0.40 and LA4330 and LA2932 have a high median, with an
255	infected fraction of 0.70 or 0.73 respectively.
256	With Fusarium we also see differences between the infected fraction of each population.
257	Interestingly LA3111 is in this case the most infected population (mean: 0.72, median: 082)
258	whereas LA4107 is the least susceptible (mean = 0.28, median = 0.11).
259	Finally, for <i>P infestans</i> , the infected fractions again show a different pattern. The data show a
260	larger spread as can be seen by the larger distance between the $1^{st}$ and $3^{rd}$ quartile and the
261	lowest and highest mean and median fraction were closer together ranging from 0.30 and 0.21
262	for LA3111 to 0.60 and 0.70 for LA4330 (Fig 2C). LA3111, one population that seems
263	particularly resistant against Alternaria and Phytophthora seems to be the most susceptible to
264	Fusarium.
265	
266	To test the significance of the differences and the effect of the different populations on infection,
267	we constructed a general linearised mixed model (glmm). We assigned experimental
268	parameters (data, box and leaf number) as random effects and tested whether there were
269	significant differences between the populations for each infecting species by looking at the
270	infection counts (y) per leaf. These models show that indeed there are highly significant
271	differences (p<0.00001) in infection rates between some populations for all three pathogens
272	tested (S Data 2).

273

#### 274 Pairwise comparisons reveal individual differences between different pathogens

275 To further determine which populations are different from each other, we performed pairwise 276 comparisons using a variant of Tukey's Honest Significant Difference test. The observed 277 pairwise differences are clearly distinct between the three pathogens. Figure 3 shows a 278 summary heatmap of the differences, with corresponding estimates for each comparison. Cells 279 with significant differences (p < 0.001) highlighted in green. All pairwise differences with their 280 95% confidence intervals are plotted in S. figure 3. Of the 63 pairwise comparisons, 32 show a 281 significant difference in infection ratio. Overall, there are more significant differences between 282 populations when it comes to Fusarium infection (15) than to Alternaria infection (10) or 283 Phytophthora (7). Interestingly, some populations show the same result for all pathogens: there 284 are no differences between LA1963 and LA2931 (both central) nor for LA2931 and LA4107 285 (south coast and central) or LA4107 and LA4117 (south coast and south mountain). Also, 286 LA1963 is always more susceptible than LA2932 and LA4117 is always more susceptible than 287 LA4330. In some cases a population in a pair is more resistant to one pathogen and more 288 susceptible to another. LA4330 is more resistant than LA3111 to Fusarium, but less resistant to 289 Alternaria and Phytophthora

290

291

#### 292 A mix of climatic and geographic variables affect pathogen resistance

293 To see whether a change in certain geographic and climatic conditions can be linked to an 294 increase or decrease of resistance rates between populations, we built new glmm using such 295 data. First we made a simple model for *Alternaria*, testing the infection counts (y) against either 296 latitude or longitude, a combination of both or an interaction of both. This showed that both 297 latitude and longitude have a significant effect (p < 0.001). A model with both parameters shows 298 a better fit, whereas a model with an interaction does not. We extended the model to include 299 both parameters (longitude + latitude) and to fit various environmental parameters (Table 1, S. 300 Data 2). We obtained the best AIC (2641.8) for a model containing altitude, annual precipitation,

301 the temperature in the wettest and the temperature in the coldest guarter. Additions of other 302 climatic data did not yield an improvement of the model. Table 1 shows that of all effects, 303 longitude is the strongest effect, followed by the mean minimum temperature in winter, the 304 annual precipitation and altitude. It should be noted that models that only take temperature 305 effects into account do not account for significance. A glmm with the infection counts set against 306 the previously identified genetic groups ( $y \sim \text{group}$ ), yields a high AIC (2705). The model with 307 the populations yields an as good AIC as the one with all available variables. This suggests that 308 no single variable has a strong, exclusive correlation to infection rate and that each population 309 represents its own micro environment with specific geographic and climate parameters that are 310 all of influence. 311 Similar to Alternaria, we tested all variables for Phytophthora and Fusarium. The pattern seen

for *Phytophthora* is almost identical to that of *Alternaria*. The AIC values are generally lower, but the trends are the same. Interestingly, *Fusarium* shows a slightly different picture. Whereas longitude is still the strongest effect, its significance is lower and the temperature in the coldest quarter of the year has a relatively large effect. The effect of altitude is not significant and differences in annual precipitation have a nearly negligible effect as well. As with *Alternaria*, the model testing for the group effect shows a lesser fit than the model per population (results for selected models can be found in S data 2).

319

#### 321 Discussion

322

323 The wild tomato Solanum chilense grows in a variety of habitats in Chile and Peru, ranging from 324 lower coastal areas to very high altitudes (>3000m). These populations experience considerable 325 variation in geographic parameters like precipitation and temperatures. It is known that S. 326 chilense has a clear demographic pattern and signs of adaptations to climatic differences 327 between different populations (Fischer et al. 2011, 2013, Nosenko et al. 2016). A demographic 328 pattern of North-South colonisation is observable with larger and more diverse populations in 329 the north of the range and smaller and less diverse populations in the south. In addition, there is 330 little to no genetic exchange between some of the southern most populations that are separated 331 by the extremely dry Atacama desert. This lead to the conclusion that S. chilense can be divided 332 in a northern, a central and two southern genotype groups (Böndel et al. 2015).

333 We hypothesised that pathogen pressures must differ a lot between such diverse geographical 334 locations and as such S. chilense should show signs of pathogen adaptations between the 335 different populations. To test our hypothesis we performed infection assays with three global 336 Solanum pathogens and with selected S. chilense populations. We observe clear differences 337 between the infection success rates of the 3 pathogens on the different S. chilense populations, 338 indeed suggesting local pathogen adaptations. We could observe a clear separation between 339 the genotype groups, only for Alternaria infection, where the central populations are more 340 susceptible than those from groups in the south. With the other pathogens, within-group 341 differences exist. Pairwise comparisons confirmed that outcomes differ within groups and 342 between pathogens. For example, a pair that shows significant differences for *Phytophthora* and 343 Alternaria infection (LA1963-LA4330) does not show this for Fusarium or the other way around 344 and very strong pairwise differences can even be seen within the previously identified genotype 345 groups (e.g. LA2932-LA4107 with *Fusarium*). We also showed that there are no generally more 346 resistant or more susceptible population. For example LA3111 is particularly resistant against

Peer.

347 *Fusarium*, but the most susceptible to *Phytophthora* and *Alternaria*.

348

349 We used a glmm to test which factors might contribute to these differences. Interestingly, 350 whereas the species as a whole, shows a strong north-south demography, our analyses show 351 that not latitude, but longitude is a very strong effect. This could at the one hand be explained 352 due to the absence of the northern most group in our analysis, but a more likely explanation is 353 the bigger geographic and associated climatic difference in the east-west gradient of the 354 species, with low altitude coastal areas in the west, and high mountains in the east. 355 Temperature differences can have large effects on the prevalence of pathogen populations as 356 shown for wild plant-pathosystems (Laine 2008) and also on crops, pathogens show adaptation 357 to different temperature regimes (Mboup et al. 2012, Stefansson et al. 2013) The mountainous 358 areas in our study have particularly cold winters and fairly low mean temperatures in summer, 359 which could be detrimental for pathogen survival or slow its growth and thus reduce pathogen 360 pressure. 361 Our results show indeed that temperature in winter as well as temperature in the wettest guarter 362 have a significant effect on infection rate. The importance of overwintering inoculum has 363 previously been shown to be a main predictor for *Podosphaera plantaginis* epidemics on 364 *Plantago lanceolata* in the next growing season (Soubeyrand et al. 2009). However, it must be 365 noted that models that only incorporate winter temperature or indeed any other single climatic 366 variables effects did not show any significance. This is in line with a between species 367 comparison for wild potato (Spooner et al. 2009) and might be related to the fact that some 368 higher altitude locations also have the highest annual precipitation rates. For example for P. 369 infestans a relative high humidity has large effects on successful sporulation (Harrison and

371

370

Lowe 1989)

372 Our climate data were extracted from worldclim.org and might not provide the whole picture. For

### Manuscript to be reviewed

373 example, precipitation data might be accurate, but do not take into account a common sea-fog 374 phenomenon, that can be observed along the coast of Chile and Peru (Cereceda and 375 Schemenauer 1991, Schemenauer and Cereceda 1992) This fog increases the local humidity 376 for several hours up to several days in certain "fog basins". Similarly, no data is available on any 377 nearby streams, rivers or irrigation canals for any of the populations. For some populations, a 378 note is available for the state of the site at the time of collection (e.g. "dry quebrada"), but it 379 remains unknown whether these features are a constant or changed in the time before 380 collection.

381

382 The best fitted models incorporate five climatic and geographic variables. Adding more variables 383 did not improve the model, mainly due to the correlations between the available climate data. 384 The strongest effects were observed for combinations of longitude and latitude together with 385 climatic variables, indicating that one or two variables alone do not determine pathogen 386 resistance. The latitude effect, which can be observed in the evolution of the species as a 387 whole, seems to be less strong in our analyses, where longitude plays a larger role. Overall, our 388 results indicate that indeed S. chilense shows local variations, which are possibly the result of 389 adaptations to local pathogen pressures. The mosaic like structure of our results indicate that 390 these resistances are likely caused by a multitude of factors. These findings are in line with 391 several inter species studies in wild potato, where no correlation could be found between geographical location of the species and resistance against P. infestans (Khiutti (2). 2015) or 392 393 A. solani (Jansky et al. 2008) To further unravel the combination of factors contributing to local 394 variations, new sampling excursions would be required, that not just collect plant and pathogen, 395 but also measure local geographic and climatic parameters.

396

In this study, several mechanisms theoretically proposed to generate stable long term
 polymorphism at host resistance and pathogen infectivity loci are shown to originate from the

399 environmental heterogeneity across populations. We conclude indeed on 1) the existence of 400 possible GxGxE interactions for given host-pathogen interactions, 2) heterogeneity in disease 401 incidence and prevalence across habitats, and most interestingly 3) a geographic mosaic of 402 exposure to different pathogens species. The presence-absence of different above- and below-403 ground pathogens on the same plants may a key component of wild systems generating 404 scenarios such as co-infection (Susi et al. 2015), cross-immunity or facilitation (Tack et al. 405 2015), with consequences for the genomics of pathogens (McMullan et al. 2015). Our research 406 did not yet focus on any genetic differences underlying the variation in infection rates and linking 407 phenotype to genotype should be one of the follow-up projects. Identification of the genes 408 involved in these resistance variations could also help to identify which plant defence 409 mechanisms are affected between populations and if there are indeed evolutionary differences 410 between defence pathways in nonhost resistance compared to resistance variation within or in 411 closely related species (Schulze-Lefert and Panstruga 2011, Stam et al. 2014). We have 412 recently shown that targeted resequencing of genes of interest can be a potent tool to calculate 413 evolutionary parameters of gene families of interest in wild tomato (Stam et al. 2016). Such 414 resequencing studies could thus help to pinpoint how molecular mechanisms are affected by 415 different pathogens as well as climatic variables.

416

#### 417 **Conclusions**

Differences in pathogen disease resistance have been well described between many wild crop relatives. Here we presented a phenotypic study that shows specific pathogen adaptations between populations of one wild tomato species *S. chilense*. We showed that there are clear differences between individual populations. Using generalised linear mixed models, we show that this variation does not follow a simple geographical cline, that multiple climatic factors are needed to explain parts of the variation and that even within previously identified genotype groups resistance properties can differ dramatically. Our study confirms a mosaic pattern in

- 425 resistance properties within one species and can form the starting point for studies unravelling
- 426 environmental effects on said properties as well as the genetic and molecular mechanisms
- 427 involved in plant-pathogen coevolution.

#### 429 Acknowledgements

- 430 We'd like to thank Hannes Rief and Giulia Schiavoni for help with the scoring of the infection
- 431 assays, Hans Hausladen, Michael Heß and Regine Ditteman (TUM Phytopathology for
- 432 providing the Alternaria strains and help with Fusarium isolation) and Brian Harrower (The
- 433 James Hutton Institute, Dundee, UK) for providing *Phytophthora*. R. Stam is supported by the
- 434 Alexander von Humboldt foundation.
- 435

#### 436 **References**

437

Agrios, G. N. 2005. Plant Pathology. - Elsevier.

- Böndel, K. B. et al. 2015. North–South Colonization Associated with Local Adaptation of the Wild Tomato Species Solanum chilense. Mol. Biol. Evol.: msv166.
- Brown, J. K. M. and Tellier, A. 2011. Plant-parasite coevolution: bridging the gap between genetics and ecology. Annu. Rev. Phytopathol. 49: 345–67.
- Cereceda, P. and Schemenauer, R. S. 1991. The Occurrence of Fog in Chile. J. Appl. Meteorol. 30: 1097–1105.
- Chaerani, R. and Voorrips, R. E. 2006. Tomato early blight (Alternaria solani): the pathogen, genetics, and breeding for resistance. J. Gen. Plant Pathol. 72: 335–347.
- Chaerani, R. et al. 2007. Assessment of early blight (Alternaria solani. J. Gen. Plant Pathol. 73: 96–103.
- Dean, R. et al. 2012. The Top 10 fungal pathogens in molecular plant pathology. Mol. Plant Pathol. 13: 414–430.
- Fischer, I. et al. 2011. Adaptation to drought in two wild tomato species: the evolution of the Asr gene family. New Phytol. 190: 1032–1044.
- Fischer, I. et al. 2013. Sequence Evolution and Expression Regulation of Stress-Responsive Genes in Natural Populations of Wild Tomato. - PLOS ONE 8: e78182.
- Forde, S. E. et al. 2004. Adaptation varies through space and time in a coevolving hostparasitoid interaction. - Nature 431: 841–844.
- Foster, S. J. et al. 2009. Rpi-vnt1.1, a Tm-22 homolog from Solanum venturii, confers resistance to potato late blight. Mol Plant Microbe Interact 22: 589–600.
- Gandon, S. and Michalakis, Y. 2002. Local adaptation, evolutionary potential and host-parasite coevolution: interactions between migration, mutation, population size and generation time. - J. Evol. Biol. 15: 451–462.
- Gavrilets, S. and Michalakis, Y. 2008. Effects of Environmental Heterogeneity on Victim– Exploiter Coevolution. - Evolution 62: 3100–3116.
- Griffiths, P. D. and Scott, J. W. 2001. Inheritance and Linkage of Tomato Mottle Virus Resistance Genes Derived from Lycopersicon chilense Accession LA 1932. - J. Am. Soc. Hortic. Sci. 126: 462–467.
- Harrison, J. G. and Lowe, R. 1989. Effects of humidity and air speed on sporulation of Phytophthora infestans on potato leaves. Plant Pathol. 38: 585–591.

Haverkort, A. J. et al. 2009. Applied Biotechnology to Combat Late Blight in Potato Caused by

Phytophthora Infestans. - Potato Res. 52: 249-264.

- Houterman, P. M. et al. 2008. Suppression of Plant Resistance Gene-Based Immunity by a Fungal Effector. PLoS Pathog 4: e1000061.
- Jansky, S. H. et al. 2008. A test of taxonomic predictivity: Resistance to early blight in wild relatives of cultivated potato. Phytopathology 98: 680–687.
- Ji, Y. et al. 2007. Sources of Resistance, Inheritance, and Location of Genetic Loci Conferring Resistance to Members of the Tomato-Infecting Begomoviruses. - In: Czosnek, H. (ed), Tomato Yellow Leaf Curl Virus Disease. Springer Netherlands, pp. 343–362.
- Jousimo, J. et al. 2014. Ecological and evolutionary effects of fragmentation on infectious disease dynamics. Science 344: 1289–1293.
- Kamoun, S. et al. 2015. The Top 10 oomycete pathogens in molecular plant pathology. Mol. Plant Pathol. 16: 413–434.
- Kavroulakis, N. et al. 2007. Role of ethylene in the protection of tomato plants against soil-borne fungal pathogens conferred by an endophytic Fusarium solani strain. J Exp Bot 58: 3853–3864.
- Khiutti, A. et al. 2015. Testing Taxonomic Predictivity of Foliar and Tuber Resistance to Phytophthora infestans in Wild Relatives of Potato. - Phytopathology 105: 1198–1205.
- Kumar, S. et al. 2013. Rapid detection and quantification of Alternaria solani in tomato. Sci. Hortic. 151: 184–189.
- Laine, A. L. 2005. Spatial scale of local adaptation in a plant-pathogen metapopulation. J. Evol. Biol. 18: 930–938.
- Laine, A.-L. 2006. Evolution of host resistance: looking for coevolutionary hotspots at small spatial scales. Proc. Biol. Sci. 273: 267–273.
- Laine, A.-L. 2008. Temperature-mediated patterns of local adaptation in a natural plantpathogen metapopulation. - Ecol. Lett. 11: 327–337.
- Lopez-Pascua, L. D. C. et al. 2010. Antagonistic coevolution across productivity gradients: an experimental test of the effects of dispersal. J. Evol. Biol. 23: 207–211.
- Ma, L. et al. 2013. A nuclear localization for Avr2 from Fusarium oxysporum is required to activate the tomato resistance protein I-2. Front. Plant Sci. 4: 94.
- Mboup, M. et al. 2012. Genetic structure and local adaptation of European wheat yellow rust populations: the role of temperature-specific adaptation. Evol. Appl. 5: 341–352.
- McMullan, M. et al. 2015. Evidence for suppression of immunity as a driver for genomic introgressions and host range expansion in races of Albugo candida, a generalist parasite.

- eLife in press.

- Michielse, C. B. and Rep, M. 2009. Pathogen profile update: Fusarium oxysporum. Mol. Plant Pathol. 10: 311–324.
- Moreno-Gamez, S. et al. 2013. Effect of disease prevalence and spatial heterogeneity on polymorphism maintenance in host-parasite interactions. Plant Pathol. 62: 133–141.
- Nosenko, T. et al. 2016. Adaptation to low temperatures in the wild tomato species Solanum chilense. Mol. Ecol. 25: 2853–2869.
- Nowicki, M. et al. 2011. Potato and Tomato Late Blight Caused by Phytophthora infestans: An Overview of Pathology and Resistance Breeding. Plant Dis. 96: 4–17.
- Peralta, I. E. et al. 2008. The taxonomy of tomatoes: a revision of wild tomatoes (Solanum section Lycopersicon) and their outgroup relatives in sections Juglandifolium and Lycopersicoides. Syst. Bot. Monogr. 84: 1–186.
- Perez, W. G. et al. 2001. Genetic Structure of Peruvian Populations of Phytophthora infestans. -Phytopathology 91: 956–965.
- Rose, L. E. et al. 2005. Natural Variation in the Pto Pathogen Resistance Gene Within Species of Wild Tomato (Lycopersicon). I. Functional Analysis of Pto Alleles. - Genetics 171: 345– 357.
- Rose, L. E. et al. 2007. Natural variation in the Pto disease resistance gene within species of wild tomato (Lycopersicon). II. Population genetics of Pto. Genetics 175: 1307–1319.
- Rose, L. E. et al. 2011. Targets of selection in a disease resistance network in wild tomatoes. -Mol. Plant Pathol. 12: 921–927.
- Salvaudon, L. et al. 2008. Genetic diversity in natural populations: a fundamental component of plant-microbe interactions. Curr. Opin. Plant Biol. 11: 135–143.
- Schemenauer, R. and Cereceda, P. 1992. Meteorological conditions at a coastal fog collection site in Peru. Atmósfera in press.
- Schulze-Lefert, P. and Panstruga, R. 2011. A molecular evolutionary concept connecting nonhost resistance, pathogen host range, and pathogen speciation. Trends Plant Sci 16: 117–125.
- Song, W. et al. 2011. Abscisic acid enhances resistance to Alternaria solani in tomato seedlings. -Plant Physiol. Biochem. PPB Société Fr. Physiol. Végétale 49: 693–700.
- Soubeyrand, S. et al. 2009. Spatiotemporal Structure of Host-Pathogen Interactions in a Metapopulation. Am. Nat. 174: 308–320.
- Spooner, D. M. et al. 2009. Tests of Taxonomic and Biogeographic Predictivity: Resistance to Disease and Insect Pests in Wild Relatives of Cultivated Potato.

- Städler, T. et al. 2005. Genealogical footprints of speciation processes in wild tomatoes: demography and evidence for historical gene flow. - Evol. Int. J. Org. Evol. 59: 1268– 1279.
- Städler, T. et al. 2008. Population genetics of speciation in two closely related wild tomatoes (Solanum section Lycopersicon). Genetics 178: 339–50.
- Stam, R. et al. 2014. The role of effectors in nonhost resistance to filamentous plant pathogens. -Front. Plant Sci. 5: 582.
- Stam, R. et al. 2016. Pooled Enrichment Sequencing Identifies Diversity and Evolutionary Pressures at NLR Resistance Genes within a Wild Tomato Population. - Genome Biol. Evol. 8: 1501–1515.
- Stefansson, T. S. et al. 2013. Local adaptation and evolutionary potential along a temperature gradient in the fungal pathogen Rhynchosporium commune. Evol. Appl. 6: 524–534.
- Susi, H. et al. 2015. Co-infection alters population dynamics of infectious disease. Nat. Commun. 6: 5975.
- Tabaeizadeh, Z. et al. 1999. Transgenic tomato plants expressing a Lycopersicon chilense gene demonstrate improved resistance to Verticillium dahliae race 2. Plant Cell Rep. 19: 197–202.
- Tack, A. J. M. et al. 2015. Below-ground abiotic and biotic heterogeneity shapes above-ground infection outcomes and spatial divergence in a host-parasite interaction. - New Phytol. 207: 1159–1169.
- Tellier, A. and Brown, J. K. M. 2007. Stability of genetic polymorphism in host-parasite interactions. Proc. Biol. Sci. 274: 809–17.
- Tellier, A. and Brown, J. K. M. 2009. The Influence of Perenniality and Seed Banks on Polymorphism in Plant-Parasite Interactions. Am. Nat. 174: 769–779.
- Tellier, A. and Brown, J. K. M. 2011. Spatial heterogeneity, frequency-dependent selection and polymorphism in host-parasite interactions. BMC Evol. Biol. 11: 319.
- Tellier, A. et al. 2011. Fitness effects of derived deleterious mutations in four closely related wild tomato species with spatial structure. Heredity 107: 189–199.

Thompson, J. N. 2005. The Geographic Mosaic of Coevolution. - University of Chicago Press.

- Thrall, P. H. and Burdon, J. J. 2002. Evolution of gene-for-gene systems in metapopulations: the effect of spatial scale of host and pathogen dispersal. Plant Pathol. 51: 169–184.
- Thrall, P. H. and Burdon, J. J. 2003. Evolution of virulence in a plant host-pathogen metapopulation. Science 299: 1735–1737.

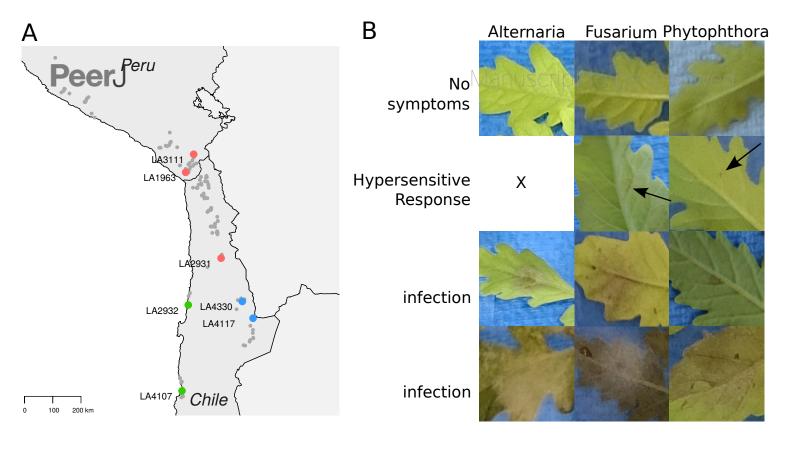
- Thrall, P. H. et al. 2001a. Short-term epidemic dynamics in the Cakile maritima–Alternaria brassicicola host–pathogen association. J. Ecol. 89: 723–735.
- Thrall, P. H. et al. 2001b. Variation in resistance and virulence among demes of a plant hostpathogen metapopulation. - J. Ecol. 89: 736–748.
- Thrall, P. H. et al. 2002a. Local adaptation in the Linum marginale-Melampsora lini hostpathogen interaction. - Evol. Int. J. Org. Evol. 56: 1340–1351.
- Thrall, P. H. et al. 2002b. Local adaptation in the Linum marginale-Melampsora lini hostpathogen interaction. - Evolution 56: 1340–1351.
- Verlaan, M. G. et al. 2013. The Tomato Yellow Leaf Curl Virus Resistance Genes Ty-1 and Ty-3 Are Allelic and Code for DFDGD-Class RNA–Dependent RNA Polymerases. - PLOS Genet 9: e1003399.
- Vogwill, T. et al. 2009. Source Populations Act as Coevolutionary Pacemakers in Experimental Selection Mosaics Containing Hotspots and Coldspots. Am. Nat. 173: E171–E176.



### Figure 1(on next page)

S. chilense populations and phenotypic observations

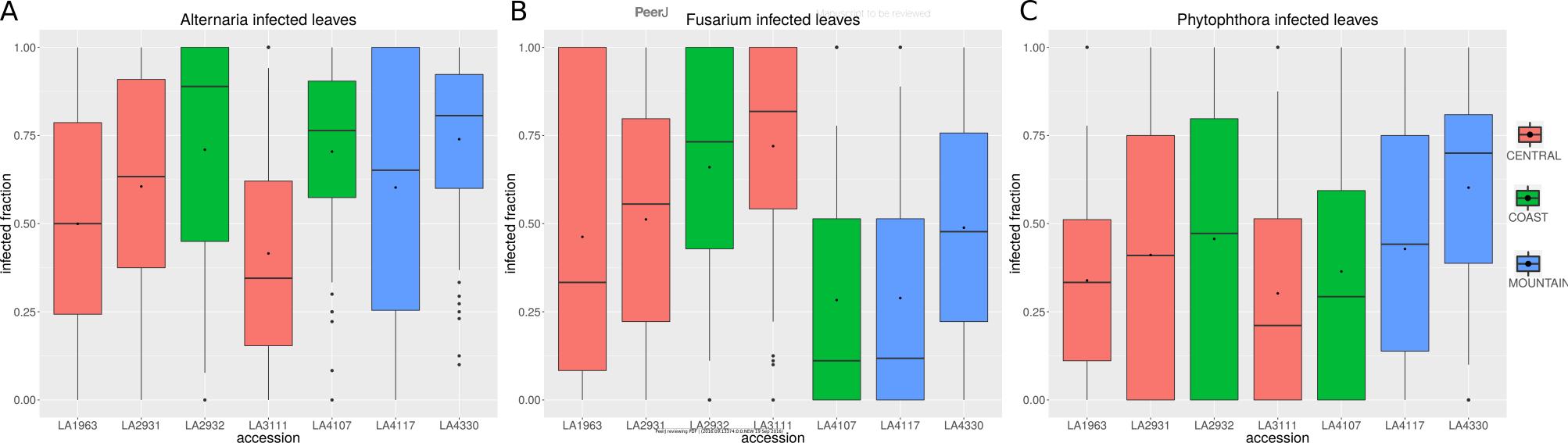
A) A map showing the populations used in this study, belonging to the central (red), southern mountainous (blue) or southern coastal (green) region. The geographic range of whole species is depicted in the background (grey dots). B) The phenotypic observations after infection range from no visible symptoms (first row) and small black necrotic lesions resembling the Hypersensitive Response (HR, second row), both scored as 'not infected', to intermediate and strong infection (third and fourth row), both scored as 'infected'. In the columns from left to right: infection with Alternaria, Fusarium and Phytophthora. We could not observe HR in the Alternaria infections.



### Figure 2(on next page)

Infected leaf fraction for different S. chilense populations

The boxplots show the median and 1st and 3rd quartile of the infected fractions per leaf for A) Alternaria, B) Fusarium and C) Phytophthora. The black dots represent the mean value for the infections. The Y axis ranges from 0 (no infected leaflets on a leaf) to 1 (all leaflets show infection). On the X axis, each population is represented. The colours correspond to the geographic regions as depicted in figure 1.



### Figure 3(on next page)

Populations with significant different infected fractions

Heatmap depicting whether a pairwise difference shows a significant result for Alternaria solani (left column), Fusarium sp. (middle column) and Phytophthera infestans (right column). Each row represents a pairwise comparison. Green cells represent a significant difference (p < 0.001 after multiple testing correction) and the numbers represent the estimated effect, with negative numbers indicating that the population mentioned on the left is less resistant than the one on the right.

Alternaria	Fusarium	Phytophthora	
-0.55183	0.10738	-0.26040	LA1963 - LA2931
-0.87557	-0.78253	-0.73009	LA1963 – LA2932
0.29545	-1.11537	0.26290	LA1963 – LA3111
-0.75 50	0.71985	-0.26015	LA1963 – LA4107
-0. +04 0-0-1	0.88139	-0.42327	LA1963 – LA4117
-1.37044	0.02266	-1.23225	LA1963 – LA4330
-0.32373	-0.88991	-0.46969	LA2931 - LA2932
0.84728	-1.22275	0.52330	LA2931 - LA3111
-0.29967	0.61247	0.00025	LA2931 - LA4107
0.08695	0.77401	0.16287	LA2931 - LA4117
-0.81860	-0.08472	-0.97185	LA2931 - LA4330
1.17102	-0.33285	0.99300	LA2932 – LA3111
0.02407	1.50237	0.46994	LA2932 – LA4107
0.41069	1.66391	0.30681	LA2932 - LA4117
0.49487	0.80518	-0.50215	LA2932 - LA4330
-1.14695	1.83522	-0.52305	LA3111 – LA4107
-0.76033	1.99676	-0.68618	LA3111 – LA4117
-1.66588	1.13803	-1.49516	LA3111 – LA4330
0.38662	0.16154	-0.16312	LA4107 – LA4117
-0.51894	-0.69719	-0.97210	LA4107 – LA4330
-0.90556	-0.85873	-0.80897	LA4117 – LA4330

### Manuscript to be reviewed

### Table 1(on next page)

Table 1

Summary of GLMM results

1 .

### Manuscript to be reviewed

Model	Alternaria	Fusarium	Phytophthora
1 y~accession	2641.8	2307.6	1893.3
2 y~Lat	2708.6	2431.3	1958.3
3 y~Long	2815.1	2490.8	1965.8
4 y~Long+Lat	2703.9	2420.6	1945.4
5 y~Long*Lat	2705.8	2419.1	1947.4
6 y~Long+Lat+Alt+AnnPrecip+TempA+TempB	2641.8	2307.6	1893.3
7 y~Altitude	2843.3	2503.5	1985.1
8 y~Temp	2843.5	2506.5	1984.1
9 y~AnnPrecip	2757.2	2457.1	1968.4
10 y~group	2705.0	2489.7	1930.2