

The quasi-universality of nestedness in the structure of quantitative plant-parasite interactions

Benoît Moury, Jean-Marc Audergon, Sylvie Baudracco-Arnas, Safa Ben Krima, François Bertrand, Nathalie Boissot, Mireille Buisson, Valérie Caffier, Melissa Cantet, Sylvia Chanéac, et al.

▶ To cite this version:

Benoît Moury, Jean-Marc Audergon, Sylvie Baudracco-Arnas, Safa Ben Krima, François Bertrand, et al.. The quasi-universality of nestedness in the structure of quantitative plant-parasite interactions. 2021. hal-03169796

HAL Id: hal-03169796 https://hal.inrae.fr/hal-03169796v1

Preprint submitted on 23 Jun 2021

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



The quasi-universality of nestedness in the structure of quantitative plant-

parasite interactions

1

2

3

10

- 4 Moury Benoît¹, Audergon Jean-Marc², Baudracco-Arnas Sylvie³, Ben Krima Safa⁴, Bertrand François⁵,
- 5 Boissot Nathalie², Buisson Mireille⁶, Caffier Valérie⁷, Cantet Mélissa^{2*}, Chanéac Sylvia⁸, Constant
- 6 Carole⁹, Delmotte François¹⁰, Dogimont Catherine², Doumayrou Juliette^{1§}, Fabre Frédéric¹⁰, Fournet
- 7 Sylvain¹¹, Grimault Valérie¹², Jaunet Thierry¹³, Justafré Isabelle¹⁴, Lefebvre Véronique², Losdat Denis¹⁵,
- 8 Marcel Thierry⁴, Montarry Josselin¹¹, Morris Cindy E.¹, Omrani Mariem^{1,2}, Paineau Manon¹⁰, Perrot
- 9 Sophie¹², Pilet-Nayel Marie-Laure¹¹, Ruellan Youna²
- ¹Pathologie Végétale, INRAE, 84140 Montfavet, France
- ²GAFL, INRAE, 84140, Montfavet, France
- 13 ³Laboratoires ASL, 755 chemin de Meinajaries, 84140 Montfavet, France
- ⁴University of Paris-Saclay, INRAE, AgroParisTech, UMR BIOGER, 78850 Thiverval-Grignon, France
- 15 ⁵Bayer Seeds SAS, Chemin de Roquemartine Mas Lamy, 13670, Saint-Andiol, France
- 16 ⁶GAUTIER SEMENCES, Route d'Avignon, 13630 Eyragues, France
- ⁷Univ Angers, Institut Agro, INRAE, IRHS, SFR QUASAV, 49000 Angers, France
- 18 *TAKII FRANCE SAS, 660 Chemin de la Crau, 13630 EYRAGUES, France.
- ⁹Sakata Vegetables Europe, Domaine de Sablas, rue du moulin, 30620 Uchaud, France
- 20 ¹⁰SAVE, INRAE, Bordeaux Sciences Agro, ISVV, 33140 Villenave d'Ornon, France
- 21 ¹¹IGEPP, INRAE, Institut Agro, Univ. Rennes, 35653 Le Rheu, France
- ¹²GEVES, 25 rue Georges Morel, CS 900024, 49071 Beaucouzé, France
- ¹³HM.Clause, 1 chemin du Moulin des Ronzières, 49800 La Bohalle, France
- 24 ¹⁴Vilmorin, Mas Pazac, 30210 Ledenon, France
- 25 ¹⁵RIJK ZWAAN France, La Vernède, 30390 Aramon, France
- 27 *present address: Bayer Seeds SAS, Chemin de Roquemartine Mas Lamy, 13670, Saint-Andiol, France
- 28 §present address: Syngenta Seeds B.V., Westeinde 62, P.O. box 2, Enkhuizen 1600 AA, The
- 29 Netherlands

26

30

Abstract

31

32

33

34

35

36

37

38

39 40

41

42

43

44

45

46 47

48

49

50 51

52

53

54

55

56

Understanding the relationships between host range and pathogenicity for parasites, and between the efficiency and scope of immunity for hosts are essential to implement efficient disease control strategies. In the case of plant parasites, most studies have focused on describing qualitative interactions and a variety of genetic and evolutionary models has been proposed in this context. Although plant quantitative resistance benefits from advantages in terms of durability, we presently lack models that account for quantitative interactions between plants and their parasites and the evolution of these interactions. Nestedness and modularity are important features to unravel the overall structure of host-parasite interaction matrices. Here, we analysed these two features on 32 matrices of quantitative pathogenicity trait data gathered from 15 plant-parasite pathosystems consisting of either annual or perennial plants along with fungi or oomycetes, bacteria, nematodes, insects and viruses. The performance of several nestedness and modularity algorithms was evaluated through a simulation approach, which helped interpretation of the results. We observed significant modularity in only six of the 32 matrices, with two or three modules detected. For three of these matrices, modules could be related to resistance quantitative trait loci present in the host. In contrast, we found high and significant nestedness in 30 of the 32 matrices. Nestedness was linked to other properties of plant-parasite interactions. First, pathogenicity trait values were explained in majority by a parasite strain effect and a plant accession effect, with no parasite-plant interaction term. Second, correlations between the efficiency and scope of the resistance of plant genotypes, and between the host range breadth and pathogenicity level of parasite strains were overall positive. This latter result questions the efficiency of strategies based on the deployment of several genetically-differentiated cultivars of a given crop species in the case of quantitative plant immunity.

Keywords

Plant resistance, pathogenicity, plant parasite, bipartite network, nestedness, modularity

58

59

60

61 62

63

64

65

66

67

68

69

70

71 72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

87 88

89

90

strains) that are targeted by this resistance.

Introduction The effectiveness of strategies of disease control based on host immunity depends on the underlying capabilities of hosts to resist infection, of parasites to overcome this resistance and on the potential of these traits to evolve. Parasites and hosts can be specialists or generalists in, respectively, their capacity to infect and their immunity. Confronting multiple genotypes of a parasite with multiple genotypes of a host reveals their interaction patterns, i.e. the magnitude and arrangement of their mutual specialization or generalism, which gives insights into the underlying genetic bases of these characters and allows implementing strategies of disease management based on host diversification. Importantly, the word "interaction" has different meanings in this context. In ecology, interactions between hosts and parasites are the effects that each of these two categories of living organisms have on each other. These host-parasite interactions can involve molecular interactions, which are attractive or repulsive forces between molecules, for example between parasite elicitors or effectors and host receptors. Finally, quantitative pathogenicity traits can be analysed thanks to statistical models that include, or not, a significant interaction between variables representing hosts and parasites. In the latter acception, "interaction" means that the model departs significantly from a purely additive model, including only a parasite effect and a host effect. Statistical interactions are used in the context of quantitative data and linear regression models, but not for qualitative binary data. The structure of any host-parasite interaction can be represented as a matrix where columns correspond to host genotypes (either inbred lines, clones or F₁ hybrids) and rows to parasite strains (either isolates, clones or populations depending on the considered parasite). Each cell in the matrix indicates the result of the pairwise confrontation between the corresponding host genotype and parasite strain. Qualitative interactions generate binary matrices with 1 and 0 grades, which correspond to successful and unsuccessful infections. Nestedness and modularity are two quantitative properties that reveal non-random distributions of 1 and 0 grades in such matrices (Weitz et al. 2013). Nestedness measures the tendency of the hosts of a parasite to have a hierarchical organization, where the set of hosts of a given parasite (a species or a genotype) is a subset (respectively superset) of that of the parasites of broader (respectively narrower) host ranges. Here, the breadth of the host range of a given parasite is defined as the percentage of host species (or genotypes) that are infected by this parasite. The same tendency is observed for host immunity (Fig. 1A): the set of parasites that are controlled by the immunity of a given host is a subset (respectively superset) of that of hosts with broader (respectively narrower) scopes of resistance. Here, the scope of the resistance of a given host is defined as the percentage of parasite species (or

92

93

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

110

111

112

113

114

115

116

117

118

119120

121

122

123

124

Modularity measures the strength by which the matrix can be divided into several modules grouping subsets of hosts and parasites characterized by successful infections, infections being rare for hosts and parasites belonging to different modules (Fig. 1B). Depending on the genetic, evolutionary and mechanistic patterns of host-parasite interactions, contrasted scores for nestedness and modularity are expected. Three main models of host-parasite interactions have been proposed for qualitative plant-parasite interactions (Fig. 1C to F; see Thrall et al. 2016 for details). These models represent the mutual specialization of hosts and parasites in terms of underlying molecular mechanisms, genetic determinism and coevolution pattern. Each one generates a specific structural pattern in the corresponding interaction matrix. Historically, the first model was the gene-for-gene (GFG) model proposed to describe interactions between crop plants and their parasites, based on genetic studies of flax and rust (Flor 1956). In this model, plant immunity is inducible and requires recognition of the parasite by its host. Recognition occurs between a host receptor and a parasite elicitor, each of them being encoded by a single gene. The loss or alteration of the elicitor in the parasite or the absence of a cognate resistance allele in its host results in infection. Here, the word 'elicitor' is used in the broad sense of a parasite component triggering plant defenses, and thus includes effectors and avirulence factors (Bent and Mackey 2007). This model is coherent with dominant resistance that involves plant proteins containing nucleotide-binding and leucine-rich-repeat domains as receptors, and that mounts hypersensitive reactions (programmed cell death) upon recognition of various kinds of parasite elicitors. In this system, a parasite strain may have universal infectivity, i.e. may be able to infect all host genotypes, if it lacks all the elicitors that correspond to the host resistance factors. Accordingly, the matrix has a global nested pattern, with partial or complete overlap of the host ranges of the parasite strains and of the resistance spectra of the host genotypes (Fig. 1C,D). Secondly, the matching-allele (MA) model was proposed to describe the self/non-self recognition system of invertebrate immunity (Grosberg and Hart 2000). In that case, infectivity requires a specific match between the host genotype and the parasite strain and, accordingly, universal infectivity is impossible. The corresponding host-parasite matrix has a modular structure. Cross-infections are frequent between hosts and parasites belonging to the same module but rare between hosts and parasites belonging to distinct modules. In extreme cases of specialization, modules can be as small as a single host-parasite pair (Fig. 1E). Mechanistically, this model is coherent with recessive plant resistance to viruses mediated by eukaryotic translation initiation factors (e.g. Sacristán and García-Arenal 2008) and with necrotrophic fungi which secrete elicitors of programmed cell death that increase plant susceptibility by allowing the fungus to feed on dying cells (Peters et al. 2019). In the context of plant necrotrophic parasites, this model is also confusingly named 'inverse gene-for-gene'

126

127

128

129

130131

132

133

134

135

136137

138

139

140

141

142

143144

145

146

147

148

149

150

151

152

153

154

155156

157

(Peters et al. 2019). Thirdly, the inverse-matching-allele (IMA) model was proposed to reflect the adaptive immune system of vertebrates, where the host resists through recognition of the parasite and infections occur when the parasite mismatches the host (Kidner and Moritz 2013; Thrall et al. 2016). The IMA model was defined in the context of multi-allelic series of resistance and pathogenicity genes. Mechanistically very similar to the GFG model, it assumes that recognition between host and parasite genotypes is highly specific. The corresponding host-parasite matrix is therefore similar to the matching-allele model but with 0 and 1 grades replaced by 1 and 0 grades, respectively (Fig. 1F). Hence, a modular pattern is the expected result when immunity levels (instead of the degree of pathogenicity) are indicated in the matrix. The distinguishing feature of the genetic models described above is that they describe qualitative binary interactions, where each host-parasite pair is characterized by its compatibility or noncompatibility. Models that describe quantitative host-parasite interactions are rare and their adequacy to represent empirical data have not been extensively tested (Lambrechts 2010; Wang et al. 2018). Analysis of quantitative plant immunity has mostly been confined to the framework of quantitative genetics and QTL (quantitative trait loci) mapping. These methods usually assume that resistance is determined by the additive effect of QTLs. More complex effects (dominance, epistasis) are rarely considered (Gallois et al. 2018). Furthermore, there are few studies of quantitative genetics and QTL mapping of parasite pathogenicity traits, especially in the case of plant parasites (Wang et al. 2018). Most importantly, these few analyses were conducted either with a set of hosts confronted to a single parasite or with a set of parasites confronted to a single host. In any case, there is a clear need for new models describing quantitative host-parasite interactions while properly accounting for the variability of both partners (Lambrechts 2010; Bartoli and Roux 2017). Moreover, previous work has shown that the outcome of analysis of matrix structure is markedly impacted when quantitative interactions are considered. Quantitative data are especially influencing the significance of nestedness (Staniczenko et al. 2013). These considerations motivated us to conduct a comprehensive analysis of the nestedness and modularity of interaction matrices to deepen our knowledge in the specialization between plants and diverse parasites using quantitative data. The objectives of this work are (i) to assess the performance of available algorithms to identify nested and modular patterns in matrices of quantitative data and (ii) to determine if these patterns are specific to each pathosystem or show a general trend. In addition, our work provides a new perspective and insight into appropriate genetic and evolutionary models for representing quantitative plant-parasite interactions and for outcomes for plant resistance management.

159

160

161

162163

164

165

166

167

168169

170

171

172

173

174

175

176

177

178

179

180

181

182

183

184

185

186

187

188

189 190 **Results** We gathered 32 matrices corresponding to 15 plant-parasite pathosystems and containing quantitative pathogenicity trait values (Table 1; Fig. 2). Among the 13 parasite species included, most were fungi or oomycetes (five and four, respectively), while bacteria, nematodes, insects and viruses were represented only once. Only three pathosystems included perennial (tree) plants and all plant species were temperate-climate crops (or crops adapted to both temperate and tropical climates). Each pathosystem included a set of strains belonging to the same parasite species and a set of accessions belonging to the same plant species with four exceptions, matrices 9, 18, 19 and 26, where accessions belonged to several closely-related plant species. Among the matrices, the number of plant accessions varied from seven to 53 (median 12) and the number of parasite strains varied from six to 98 (median 11.5). The number of matrix cells varied from 49 to 1470 (median 180). For most pathosystems, we analyzed several matrices corresponding to either different pathogenicity traits, different plant-parasite sets or different experiments. In order to meet the requirements of methods that allow the estimation of nestedness and modularity of matrices, the pathogenicity traits in each matrix were standardized into integer values ranging from 0 (minimal plant resistance and/or maximal parasite pathogenicity) to 9 (maximal plant infection and/or minimal parasite pathogenicity). We then tested for the occurrence of nestedness and modularity. For significance assessment, the nestedness/modularity scores of the matrices derived from experimental data were compared to those of simulated null-model matrices that are not expected to possess any nested or modular pattern (Supplementary Methods 1). Nestedness (or modularity) is significant if the actual matrix is more nested (or modular) than at least 95% of the matrices simulated under a given null model (black numbers on grey background in Tables 2, 3 and 4). As there are many possible null models and because their choice is crucial to conclude about the significance of nestedness or modularity, we analyzed the performance of the different available nestedness/modularity algorithms and of different null models by estimating their type I and type II error rates through a simulation approach (Supplementary Methods 1; Tables S1 to S18). Ubiquitous nestedness in quantitative plant-parasite interactions First, we evaluated the performance of two algorithms, WINE and wNODF (Galeano et al. 2009; Almeida-Neto and Ulrich 2011), to estimate the nestedness of the 32 matrices. Simulations revealed that statistical significance with both null models C1 and R1 (or C2 and R2) provided the lowest false positive rates for nestedness (Supplementary Methods 1; Tables S1 and S2). Under null models C1

192

193

194

195

196

197

198

199

200

201

202

203

204

205

206

207

208

209

210

211

212

213

214

215

216

217

218

219

220

221

222

and R1, matrices are generated column by column or row by row, respectively, and the cell numbers are chosen randomly in order that (i) the marginal sums of cells and (ii) the numbers of zero-valued cells are kept the same as in the actual matrix. To generate matrices under null models C2 and R2, the cell values of the actual matrix are shuffled column by column or row by row, respectively. With the WINE algorithm, nestedness values were quite high in general (from 0.46 to 1.04; mean 0.77 on a scale varying from 0 to ≈1). Thirty of the 32 matrices showed significant nestedness (pvalues ≤ 0.05) with null models C1, R1, C2 and R2 (Table 2). Only matrices 21 and 32 were not significantly nested with either null model C1, R1, C2 or R2. With the wNODF algorithm, nestedness estimates varied from 6.1 to 75.4 (mean 38.2) on a scale varying from 0 to 100 and nestedness was significant for only 19 of the 32 matrices with null models C1, R1, C2 and R2 (Table 2). This lower number of matrices showing nestedness is consistent with the lower statistical power of wNODF compared to WINE (Supplementary Methods 1). As both methods are based on different principles, the correlation of their nestedness scores among the 32 matrices is only moderate (Pearson's r = 0.37; p-value = 0.038). Importantly, unlike the WINE method, wNODF cannot estimate the nestedness of matrices devoid of zero-valued cells and underestimates nestedness when zero-valued cells are scarce. Indeed, most of the matrices significantly nested with WINE but not significantly nested with wNODF contained few zero-valued cells, most of which being distributed on a single row or column. Consequently, the discrepancy between results obtained by WINE and wNODF may be a bias due to the lack or peculiar distribution of the zero-valued cells. Five more matrices (numbers 13, 15, 20, 29 and 31) were significantly nested with wNODF if 1-valued cells or 1- and 2-valued cells were transformed into 0-valued cells. With wNODF (but not with WINE), several matrices were less nested than at least 95% of the matrices simulated under one or several null models (white numbers on black background in Table 2), a property that we will name anti-nestedness. Matrix 14 was significantly anti-nested with null models N, C1, R1 and R2. For six other matrices, significant anti-nestedness was detected with one or a few null models, a bias attributable to the small number of 0-valued cells (matrices 12, 23 and 24) which disappeared largely when 1- and 2-valued cells were transformed into 0-valued cells. Overall, taking into account the limitations of the wNODF algorithm, our analysis revealed that the huge majority of the matrices (30/32; 94%) were significantly nested. Investigation of the biological significance of nestedness Adequacy of an additive linear regression model for pathogenicity matrices

224

225

226

227

228229

230

231

232

233

234

235

236

237

238

239

240

241

242

243244

245

246

247

248

249

250

251

252

253

254

255

The high and significant nestedness observed among most of the analysed matrices suggests that an additive model combining pathogenicity QTLs in the parasites and resistance QTLs in the hosts, with no QTL x QTL interactions between hosts and parasites, would fit well with the data (Fig. 1G). We evaluated the performance of the linear regression model: 'pathogenicity' ~ 'parasite strain' + 'plant accession', with no interaction term, on the datasets. For each plant accession-pathogen strain pair, the mean pathogenicity value was considered for the 'pathogenicity' variable. The 'parasite strain' and 'plant accession' effects were highly significant (p-value < 0.0012), except for matrices 21 and 32 which were the only ones not significantly nested according to the WINE method (Table 2). Omitting these two matrices, the multiple coefficient of determination (R2) indicating model fit varied from 0.49 to 0.98 (mean 0.75) (Table 1), which lends support to the suggested genetic model. Moreover, the multiple R² values of the linear regression model were significantly correlated with the nestedness scores obtained with the WINE algorithm (Pearson's r = 0.73; p-value = 2.6e-06) across the 32 matrices. They were only marginally correlated with the nestedness scores of the wNODF algorithm (r = 0.32; p-value = 0.07). Evaluating potential trade-offs: Host range breadth vs. pathogenicity in parasites and scope vs. efficiency of resistance in host plants The ubiquitous nestedness detected suggests a positive correlation between the host range breadth, i.e. the percentage of host accessions that a parasite can efficiently infect, and the pathogenicity level of the parasite. Similarly, a positive correlation is expected between the scope of the resistance and the resistance efficiency of the plants. Given the continuous distribution of the quantitative pathogenicity traits, we defined arbitrary pathogenicity thresholds to distinguish host and non-host accessions for a given parasite strain, and to distinguish parasite strains included or not included in the scope of the resistance of a given plant accession. Nine thresholds were defined, varying from 10% to 90% of the maximal pathogenicity value in the whole matrix by increments of 10%, and allowed estimating the percentage of plant accessions included in the host range of each parasite strain (i.e. the host range breadth) and the percentage of parasite strains included in the scope of resistance of each plant accession. The mean Pearson's coefficient of correlation (r) between host range breadth and pathogenicity varied from 0.20 to 0.38 across the different threshold values (mean 0.31). Depending on the threshold, from 23.1% (6/26 matrices) to 40.6% (13/32) (mean 31.9%) of the matrices showed significantly positive r values, whereas from 0 (0/11) to 9.7% (3/31) (mean 4.8%) of the matrices showed significantly negative r values (Fig. 3). Note that the coefficient of correlation could not be calculated for several matrices for some of the thresholds because of the

257

258

259

260

261

262

263

264

265

266

267

268

269

270

271

272

273

274

275

276

277

278

279

280

281

282

283

284

285

286287

lack of pathogenicity values above (for correlation between host range breadth and pathogenicity) or below (for correlation between resistance scope and efficiency) that threshold. The mean r between resistance scope and efficiency varied from 0.18 to 0.59 across the different threshold values (mean 0.39). Depending on the threshold, from 25.0% (6/24) to 46.9% (15/32) (mean 35.8%) of the matrices showed significantly positive r values, whereas from 0 (0/32) to 9.4% (3/32) (mean 2.9%) of the matrices showed significantly negative r values (Fig. 3). Rare cases of modularity in quantitative plant-parasite interactions We applied five algorithms to estimate the modularity of the 32 matrices (Newman and Girvan 2004; Clauset et al. 2004; Newman 2006; Blondel et al. 2008; Traag and Bruggeman 2009; Supplementary Methods 1). By maximizing a modularity score, these algorithms estimate the optimal number of modules and the distribution of plant and parasite genotypes in the modules. Modularity scores were low overall, with a maximum of 0.240 and a mean of 0.075, on a scale varying from 0 to 1 (Tables 3 and 4). The fast greedy, louvain and leading eigenvector methods provided highly similar modularity scores among the 32 matrices, with Pearson's coefficients of correlation r > 0.91 (p-values ≤ 1e-12). Scores of the edge betweenness method were highly correlated with the previous three methods (0.58 < r < 0.74; p-values $\le 6e-04$) whereas scores of the *spinglass* method were moderately correlated with the previous ones (0.39 < r < 0.61; p-values ≤ 0.027). Our analysis of the performance of these methods showed that the spinglass algorithm had a very low rate of false positive modularity, whatever the null model (Supplementary Methods 1). In contrast, the fast greedy, louvain and edge betweenness algorithms had high rates of false positive modularity with several null models, except models S (where cell values of the actual matrix are shuffled, with no constraints on row or column marginal sums), C2 and R2 (Tables S7-S9). According to the spinglass method, six matrices (numbers 5, 6, 10, 11, 14 and 17b) were significantly modular with a majority of null models (Table 3), though their modularity scores were low (≤ 0.102). Depending on the matrix, spinglass defined an optimal number of two or three modules, which provided the maximal modularity score (Table 3; Fig. 4). In addition, matrices 8 and 22 were only significantly modular with one null model. Evidence of modularity with the edge betweenness, fast greedy, louvain and leading eigenvector methods was scarce, significant modularity being usually observed for one of the null models S, C2 or R2 only and not for all methods (Table 4). With all modularity methods, several matrices were less modular than at least 95% of the matrices simulated under one or several null models (white on black numbers in Tables 3 and 4), a property

289

290

291

292

293

294

295

296

297

298

299

300

301

302

303

304

305

306

307

308

309

310

311

312

313

314

315

316

317

318319

that we will name anti-modularity. For spinglass, only matrix 7 was significantly anti-modular with null models N, C1 and R1 (Table 3). The other methods detected significant anti-modularity in most matrices with most null models but suffered high rates of false positive anti-modularity for many null models (Supplementary Methods 1; Tables S11-S13). Considering only matrices that are significantly anti-modular with both null models C2 and R2, which correspond to the lowest rates of false positives (null model CR2 in Tables S11 to S13), 13 matrices were significant with at least two methods and three (matrices 17, 28 and 29) were significant with the four methods (Table 4). Results obtained with the other matrices varied according to algorithms and null models, showing both significant modularity and anti-modularity, which could be due to low type I error performances of the algorithms for detection of modularity and/or anti-modularity. Investigation of the biological significance of modularity We examined the relevance of the detected modules for the six matrices showing significant modularity with most null models with spinglass (Table 2) by analysing whether the plant and parasite genotypes belonging to each module shared common properties (common resistance gene or QTL for plants; common pathogenicity factor for parasites; common origin for plants or parasites). For matrix 5 (Puccinia hordei-barley), two modules were detected (Fig. 4). The first one grouped the five accessions with resistance QTLs Rphq3 and Rphq11, showing delayed infection with most isolates of the second module, and one accession carrying QTLs Rphq1, Rphq2 and Rphq3, showing delayed infection with almost all isolates (González et al. 2012). The second module contained four accessions with either no resistance QTL or QTL Rphq18, that were quickly infected by almost all isolates. The country of origin or date of collection of the isolates did not explain their distribution in the two modules (Marcel et al., 2008). For matrix 6 (Venturia inaequalis-apple), three modules were detected. The first one grouped the eight accessions carrying QTL T1 and the four V. inaequalis isolates collected on apple trees carrying T1 (Laloi et al. 2017). The two other modules grouped (i) the remaining accessions that were either carrying no resistance QTL or QTLs F11 or F17 that have only a low effect on disease reduction and (ii) isolates collected on these accessions. One of these modules grouped a single isolate and a single accession. Infections were on average high within all modules and low between any pair of modules. Two modules were also detected for matrix 14 (Zymoseptoria tritici-bread wheat). These modules could be partially explained by the interaction between the resistance gene Stb6 (Saintenac et al. 2018), that confers a high level of resistance in the absence of a hypersensitive response, and the

321

322

323

324

325

326

327

328

329

330

331

332

333

334335

336337

338339

340

341

342

343

344

345

346

347

348

349

350

pathogen avirulence gene AvrStb6 (Zhong et al. 2017). Six of the eight cultivars in the first module carry Stb6, while at least six of the seven cultivars in the second module do not carry Stb6. Moreover, the 44 fungal isolates structuring the first module are pathogenic on Stb6 while the 54 isolates from the second module are either pathogenic or not pathogenic on Stb6. Concerning matrices 10, 11 (Podosphaera xanthii-melon) and 17b (Phytophthora capsici-pepper), three modules were detected but there was no evidence of similarity in the genetic composition of accessions, the presence of particular resistance genes or QTLs or the origin of isolates belonging to a same module. Modularity of reverse matrices To test the occurrence of IMA patterns (Fig. 1F), we also analyzed the modularity of the 32 matrices transformed such that a grade of 0 corresponds to the maximal plant susceptibility and grades 1 to 9 correspond to the range of increasing plant resistance (hereafter "reverse matrices"). Using the spinglass algorithm, four matrices (numbers 10, 11, 14 and 15) showed significant but low modularity (≤0.078) with either null models C1 and R1 or C2 and R2. Depending on the matrix, spinglass defined an optimal number of two to five modules (Fig. 5). The modules identified in reverse matrices 14 and 15 using the spinglass algorithm were biologically more meaningfull than the two modules previously identified for matrix 14. Matrices 14 and 15 correspond to two different phenotypic traits measured in the same plant-parasite interactions (i.e. necrosis and sporulation, respectively). Interestingly, modules identified in the two matrices were similar but not identical since five modules were identified in matrix 14 and four modules were identified in matrix 15. This may reflect differences in the genetic determinism of the two phenotypic traits measured or differences in the mechanisms of various Stb resistance genes. For matrix 14, three modules correspond to the presence of resistance genes Stb7 (one cultivar), Stb9 (three cultivars) and Stb6 (four cultivars), one module to cultivars carrying various Stb genes (three cultivars), and one module to susceptible (or partially resistant) cultivars (four cultivars). For matrix 15, the modules corresponding to the presence of Stb6 and Stb9 are also identified (with an additional cultivar in the Stb6 module), the module corresponding to susceptible cultivars as well (with two additional cultivars), and the cultivar Salamouni carrying Stb13 and Stb14 forms the fourth module. As above, there was no evidence of similarity in the composition of accessions and isolates belonging to the same module for reverse matrices 10 and 11.

Overall, considering both the initial and reverse matrices, our analysis revealed that only a minority of the matrices (7/32; 22%) were significantly modular.

Discussion

351

352

353

354

355356

357

358

359

360

361

362

363

364

365

366

367

368

369

370

371

372

373

374

375

376

377

378

379

380

381

382

383

There is nothing more fundamental to the concepts in Plant Pathology as a science and to the practical strategies used for managing plant health than the host range of a parasite and the scope of resistance of a plant (Morris and Moury 2019). Based on the patterns in matrices of plant-parasite interactions, we can conceive and test hypotheses about the molecular and evolutionary processes that underlie plant-parasite interactions, develop robust diagnostic tools, design breeding programs and strategies for deploying resistant cultivars, and construct models to anticipate disease emergence. Given the complexity of the mechanisms involved in disease, it would be reasonable to assume that the particularities of each pathosystem would be an impediment to identifying universal principles that can guide these efforts. However, here we have used network-based analyses to reveal the quasi-universal principle that the structure of quantitative matrices of plant-parasite interactions is nested. Indeed, evidence of nestedness was found in 94% (30/32) of the matrices that we analyzed and one of the two non-nested matrices (number 32) was one of the smallest ones, which may have precluded the detection of a significantly nested pattern. Our results were based on statistically robust analyses of quantitative assessments of compatible interactions between hosts and parasites for large interaction matrices involving from 49 to 1470 (median 180) host-parasite combinations. Quantitative data are key to the accuracy and genericity of these analytical methods. Indeed, in a study of 52 published matrices containing data on plant-pollinator, plant-seed disperser and parasitoid-host interactions, Staniczenko et al. (2013) found evidence of nestedness in only 3% of matrices including quantitative data, whereas the same matrices considered in a binary manner showed evidence of nestedness in 98% of cases. Network theory has its origins in the study of social networks and in ecology of interacting organisms (Patterson and Atmar 1986). Ecological networks are typically identified by counting in natura the interactions between (or co-occurrence of) two sets of taxa. Evidence of nestedness was frequent for all kinds of matrices, including interactions between hosts and symbionts, either mutualistic or parasitic (Bascompte et al. 2003; Joppa et al. 2010; Dormann et al. 2017). A number of factors that are external to the interacting organisms can affect properties of such ecological networks. For example, nestedness increases with the abundance of taxa (Joppa et al. 2010; Staniczenko et al. 2013; Suweis et al. 2013; Valverde et al. 2018), with heterogeneous distribution of connections, i.e. of numbers of links between interacting taxa (Jonhson et al. 2013), with the occurrence of broad

385

386

387

388

389

390

391

392

393

394

395

396

397

398

399

400

401

402

403

404

405

406

407

408

409

410

411

412413

414

415

416

connectivities (Feng and Takemoto 2014) and with spatially-limited interactions between taxa (Valverde et al. 2017). These analytical methods were recently used to analyse host-symbiont interactions resulting from cross-inoculation experiments, where every host taxon was inoculated with every symbiont taxon, and the compatibility of each host-symbiont pair was reported in the matrix (Flores et al. 2011; Flores et al. 2013; Weitz et al. 2013). The structural patterns of such matrices, where all host-symbiont pairs are evaluated under the same experimental and environmental conditions, are mainly the result of intrinsic, mostly genetic, differences between host or symbiont taxa. Network analyses can also be strongly affected by the choice of null models (Gotelli and Graves 1996). This is why we conducted a thorough evaluation of the performance of several null models with simulations (Supplementary Methods 1). The null models should keep, as much as possible, everything identical to the actual matrix apart from the pattern of interest, nestedness or modularity. Many null models have unacceptably loose constraints. For example, null models that do not force row or column marginal sums to be constant create distributions of taxa that do not match those usually observed, leading to falsely positive nestedness (Brualdi and Sanderson 1999; Joppa et al. 2010). Accordingly, high rates of false positives were observed with null models N and S in our simulations (Tables S1 and S2). Since parasites typically differ greatly in the number of hosts they exploit and the efficiency with which they exploit them, we did not want null models to detect significant nestedness when the heterogeneity of infection was shuffled randomly among hosts, as was frequently observed for null models N and S with test matrices M1R to M5R (Tables S1 and S2). Null models R1 and R2 that force row marginal sums to be constant avoided this problem (Tables S1 and S2). The same was true for the scope and efficiency of resistance that differ greatly between plant accessions. In that case, the C1 and C2 null models efficiently avoided an excess of falsely positive nestedness due to the hererogeneity of resistance (because C1 and C2 are equivalent to R1 and R2 when the rows and columns of the matrix are exchanged, which leaves the nestedness scores unchanged; data not shown). Overall, to account for both plant resistance and parasite infection heterogeneities, we found that the CR1 (or CR2) null model, that combines null models C1 and R1 (or C2 and R2, respectively), is the most efficient as it showed acceptable type I error rates (Supplementary Methods 1). Null model B, based on Patefield's (1981) algorithm, maintains both the row and column marginal sums of the actual matrix. However, it does not maintain the connectance (i.e. number of non-zero-valued cells of the matrix), which has a strong impact on the estimation of nestedness. Consequently, the type I error rates associated with null model B were frequently higher than those obtained with models CR1 or CR2. Moreover, using quantitative instead of binary data

418

419

420

421

422

423

424

425

426

427

428

429

430

431

432

433

434

435

436

437

438

439

440

441

442

443

444

445

446

447448

contributed to lowering the nestedness false positive rate (Staniczenko et al. 2013; Dormann et al. 2017). Overall, we obtained strong and consistent evidence of nestedness for almost all matrices (except matrices 21 and 32), whatever the parasite type, the plant species or the pathogenicity trait measured. Nestedness was linked to two important features of quantitative plant-parasite matrices: (i) scarcity and/or low level of statistical interactions between plant and parasite genotypes in terms of infection intensity and (ii) lack of trade-offs between host range and pathogenicity among parasite strains and between efficiency and scope of the resistance among plant accessions. The former feature is supported by the fact that an additive linear model - containing only a plant accession effect and a parasite strain effect with no interaction term - showed high multiple coefficients of determination (from 0.49 to 0.98) across matrices (Table 1). This result is compatible with a genetic model where pathogenicity in the parasite and resistance in the host plant are determined by a varying number of QTLs, but the statistical interaction between effects of QTLs from the parasite and QTLs from the host is rare and/or of small magnitude (Table 1; Fig. 1G). In other words, plants and parasites differ by their QTL assemblage (i.e. QTL numbers and/or effects) but plant resistance QTLs have similar effects towards all parasite strains and, reciprocally, parasite pathogenicity QTLs have similar effects towards all plant genotypes. Quantitative models usually used to analyse empirical data on plant-parasite interactions are quite simplistic, e.q. assuming or not a statistical interaction between plant and parasite genotypes (Parlevliet 1977). Models that are more complex have been proposed in the frame of theoretical modelling (e.g. Fenton et al. 2009) but their relevance to represent biological data was not evaluated. Importantly, we do not argue that evidence of nestedness supports a single genetic model of plant-parasite interaction. Instead, we suggest that an additive linear model with a plant accession and a parasite strain effects is the simplest model that accounts for the empirical data but we cannot exclude that other models could be suitable, like the modified GFG model of Fenton et al. (2009). A future challenge, requiring more in-depth genetic studies, would be to evaluate the adequacy of these different models to represent empirical plant-parasite interactions. New analytical methods can provide a better understanding and quantification of host-parasite genetic interactions, such as the host-parasite joint genome-wide association analysis recently developed by Wang et al. (2018). Applied to the Arabidopsis thaliana-Xanthomonas arboricola pathosystem, this model showed that 44%, 2% and 5% of the phenotypic variance could be explained respectively by the parasite strain, the host accession and the parasitehost interaction. As in our results, only a small parasite-host interaction effect was detected.

450

451

452

453

454

455

456

457

458

459

460

461

462

463

464

465

466

467

468

469

470

471

472

473

474

475

476

477

478

479

480

481

482

The latter feature of quantitative plant-parasite matrices is supported by the fact that we observed a majority of positive, rather than negative correlations (i.e. trade-offs), between the infectivity and the breadth of host range of parasites on the one hand and, especially, between the efficiency and scope of the resistance of plants on the other hand (Fig. 3). Few studies have examined the relationships between the scope and efficiency of plant resistance. In contrast with our results, Barrett et al. (2015) hypothesized evolutionary trade-offs between resistance efficiency and scope because quantitative resistance had a broader scope compared to qualitative resistance in the *Linum* marginale – Melampsora lini interactions. The difference between our studies could be that we focussed on quantitative resistance and included few qualitative resistance genes in our dataset (or these were overcome by most parasite strains). The positive correlation between parasite infectivity and host range breadth contrasts with qualitative host-parasite interactions and especially the GFG model, where the expansion of the host range of parasites is associated with a cost in fitness during infection of the previous hosts. Such so-called "virulence costs" have been experimentally measured in many plant-parasite systems, including viruses (Jenner et al. 2002; Desbiez et al. 2003; Janzac et al. 2010; Poulicard et al. 2010; Fraile et al. 2011; Ishibashi et al. 2012; Khatabi et al. 2013), fungi (Bahri et al. 2009; Huang et al. 2010; Caffier et al. 2010; Bruns et al. 2014), oomycetes (Montarry et al. 2010), bacteria (Vera Cruz et al. 2000; Leach et al. 2001; Wichmann and Bergelson 2004) or nematodes (Castagnone-Sereno et al. 2007), and could explain why universal pathogenicity is not fixed in pathogen populations (Tellier and Brown 2011). For quantitative plant resistance, few studies have estimated the occurrence of pathogenicity costs. Montarry et al. (2012) showed a cost for PVY to adapt to a quantitative pepper resistance when inoculated to a susceptible pepper genotype, whereas Delmas et al. (2016) showed, on the opposite, that there was no fitness cost associated with the adaptation of *Plasmopara viticola* to partially resistant grapevine varieties. Fournet et al. (2016) even highlighted that nematode populations that had adapted to potato quantitative resistance were more pathogenic on a susceptible potato genotype than were naïve nematode populations. The present study focused mostly on interactions between plants and parasites at the intraspecific level, but other studies have revealed a similar trend when strains of a given parasite species are confronted with numerous plant species. For example, a positive correlation was observed between species host range and pathogenicity for *Pseudomonas syringae* (Morris et al. 2000, 2019). For this bacterium, the most pathogenic strains were also the most ubiquitous in the environment, suggesting also an absence of trade-off between host range and dispersal capability or survival in the environment (Morris et al. 2010). In contrast to nestedness, we obtained little evidence of modularity among the matrices that we analysed. Modularity scores were low for all matrices. In only seven matrices, representing either

484

485

486

487

488 489

490

491

492

493

494

495

496

497

498

499

500

501

502

503

504

505

506

507

508

509

510

511

512

513

and in the following text.

infection or resistance scores (i.e. reverse matrices), did we detect significant modularity with a majority of null models (Tables 3 and 4; Fig. 4 and 5). For four of these matrices (matrices 5, 6, 14 and 15), modularity was linked to the presence of particular resistance genes or QTLs in the plant accessions and, for the parasite strains, to the presence of particular avirulence genes or to a common origin in terms of host genotype. For the remaining matrices (10, 11 and 17b), no common property could be found for plant accessions and parasite strains belonging to the same module. The lack of modularity of infection matrices and of reverse matrices suggests that the MA and IMA genetic models are either inadequate to represent the structure of quantitative plant-parasite interactions or explain only marginally their structure (Fig. 1E,F). **Conclusion** The ubiquitous nested patterns observed in quantitative plant-parasite interaction matrices have important implications for our understanding and management of plant diseases. They can help infer the underlying genetic bases of quantitative aspects of disease manifestation and their evolution. Our results are compatible with an additive model comprising a plant resistance effect, a parasite pathogenicity effect and no (or little) plant-parasite interaction effect. A major enigma that we highlight is the apparent lack of trade-off between pathogenicity and host range breadth among strains of a parasite, which has important implications on the efficiency of plant resistance management through cultivar rotation, mixtures or mosaics. Indeed, these strategies rely at least in part on a counter-selection of the most pathogenic parasite strains by a diversification of plant cultivars (Brown 2015). The efficiency of these strategies would certainly be reduced in absence of costs of adaptation to plant resistance. Therefore, in absence of such costs, the efficiency of the rotation, mixtures or mosaic strategies would rather depend on barrier effects designed to limit parasite dispersal in agricultural landscapes. **Materials and Methods Datasets** To be able to analyse plant-parasite interaction networks, we selected datasets containing at least 6 plant accessions and 6 parasite strains. A brief description of these datasets is provided in Table 1

515

516

517

518

519

520

521

522

523

524

525

526

527

528

529

530

531

532

533

534

535

536

537

538

539540

541

542

543

544

Matrices 1 to 4: Pseudomonas syringae-Prunus armeniaca (apricot) Nine strains of *Pseudomonas syringae*, the causal agent of bacterial canker of apricot, were inoculated on dormant tissues of twenty apricot cultivars chosen according to their differential susceptibility in orchard conditions. The strains were chosen mainly within phylogroups 1 and 2, the most abundant groups of *P. syringae* in contaminated apricot orchards in France (Parisi et al. 2019). Seven strains were isolated from symptomatic trees and two in crop debris and soil. Bacterial inoculum was prepared by cultivation on King's B medium for 48h at 24°C. The concentration of the bacterial suspension was adjusted at 108 CFU.ml⁻¹. A volume of 25 μl of inoculum was deposited at the level of a wound made superficially with a scalpel on the bark of one-year-old twigs grown in orchard. Five months after inoculation, twigs were removed and the length of flat zone around the inoculation point at the surface of the shoot (matrices 1 and 2) and the length of browning zone around the inoculation point below the bark of the shoot (matrices 3 and 4) were measured. Two independent tests were performed in 2017 (matrices 1 and 3) and 2018 (matrices 2 and 4). Matrix 5: Puccinia hordei-Hordeum vulgare (barley) Fourteen Puccinia hordei isolates (from Europe, Morocco, Israel and the USA) were inoculated on a differential series of 12 H. vulgare lines carrying different Rphq QTLs (González et al. 2012). The first seedling leaves of each barley line were inoculated with ≈240 spores/cm². The relative latency period (RLP) (Table 3 in González et al. 2012) was estimated by the number of hours from inoculation to the moment at which 50% of the ultimate number of uredinia was visible. Matrices 6 to 8: Venturia inaequalis-Malus domestica (apple tree) Grafted plants of different apple accessions (Malus domestica) carrying resistance QTLs (T1, F11, F17, F11 + F17 or T1 + F17 or ro ro resistance QTL were inoculated in controlled conditions with isolates of Venturia inaequalis, a fungal pathogen responsable of apple scab. The percentage of sporulating leaf area was assessed from 8 to 21 days post inoculation (dpi) on a scale with eight levels: 0 = no visible symptom, 0.5 = 0-1%, 3 = 1-5%, 7.5 = 5-10%, 17.5 = 10-25%, 37.5 = 25-50%, 62.5 = 50-75%, and 87.5 = 75–100%. Matrix 6 (Laloi et al. 2017) consisted of interactions between 10 V. inaequalis isolates sampled in one orchard (Angers, France) on apple trees carrying T1, F11+F17, T1+F11+F17 or no QTL and 14 apple accessions carrying the matching resistance QTL or no QTL.

546

547

548

549

550551

552

553

554

555

556

557

558559

560

561

562

563

564

565

566567

568

569

570

571

572

573

574

Matrix 7 (Caffier et al. 2016) consisted of interactions between 14 V. inaequalis isolates sampled in one orchard (Angers, France) on apple trees carrying or not T1 and 12 apple accessions carrying or not T1 (with six accessions for each of the two classes). Matrices 6 and 7 represent the Area Under the Disease Progress Curve (AUDPC) of the percentage of sporulating leaf area from eight to 21 dpi. Matrix 8 (Caffier et al. 2014) consisted of interactions between 24 V. inaequalis isolates sampled in two orchards (Lanxade and Villeneuve d'Ascq, France) on apple trees carrying F11, F17, F11+F17 or no QTL and eight apple accessions carrying the matching QTL or no QTL (with two accessions for each of the four classes). Matrix 8 represents the percentage of sporulating leaf area 14 dpi. Matrix 9: Botrytis cinerea-Solanum lycopersicum / Solanum pimpinellifolium (tomato) Leaves of 12 tomato accessions (six domesticated accessions of Solanum lycopersicum and six accessions of the close wild relative S. pimpinellifolium) were infected with single droplets of spore suspensions of 94 B. cinerea isolates. The size of lesions was measured from digital images 72 hours after inoculation (Soltis et al. 2019). One isolate was poorly infectious on all tomato accessions (grade 0 after data transformation) and was withdrawn. Matrices 10 and 11: Podosphaera xanthii-Cucumis melo (melon) Nineteen melon differential lines were inoculated with 26 Podosphaera xanthii isolates collected in 2013 and 2014 in melon, squash, watermelon and cucumber crops in Southern Europe or Northern Africa (France, Spain, Italy, Morocco, Turkey, Greece). Each P. xanthii isolate was propagated on cotyledons of Lagenaria ciceraria for seven days and spores were blown on eight leaf disks per melon line-P. xanthii isolate combination using an inoculation tower (Perchepied et al. 2005). Sporulation intensity was scored 14 days after inoculation and data were transformed in percentage of leaf disk surface using the class mean as suggested by Nicot et al. (2002): 0 = 0%, 1 = 2.5%, 2 = 7.5%, 3 =17.5%, 4 = 37.5%, 5 = 67.5%, 6 = 82.5%, 7 = 92.5%, 8 = 97.5%, and 9 = 100%. The mean score for melon accession - P. xanthii isolate combinations was reported in matrix 10. For matrix 11, 31 isolates were inoculated to 19 differential lines on leaves of entire plants. The sporulation intensity was scored similarly as for leaf disks using a 0 to 9 scale. The mean score for melon accession - P. xanthii isolate combinations was reported in matrix 11.

Matrices 12 to 15: Zymoseptoria tritici-Triticum aestivum (bread wheat) or T. turgidum subsp. durum

575

576

577

578

579

580

581

582

583

584

585

586

587

588

589

590

591

592

593

594

595

596

597

598

599

600

601

602

603

604

605

606

607

(durum wheat)

Matrices 12 and 13 were built by inoculating 12 lineages from a durum wheat landrace called Mahmoudi with 15 Zymoseptoria tritici isolates. The 12 plant lineages were fixed from individivuals coming from a single field at Journine in Tunisia and corresponded to 12 different multilocus genotypes (MLGs) as defined previously by Ben Krima et al. (2020). The 15 isolates were collected in situ either from the landrace Mahmoudi or from the cultivar Karim, at Journine in 2018. Matrices 14 and 15 were built by inoculating 15 bread wheat cultivars (Triticum aestivum), 12 of which carrying different Stb resistance genes, with 98 Z. tritici isolates collected mostly on cultivars Apache and Premio, all over France between 2009 and 2010. These bread wheat cultivars belong to a series of differential genotypes used to characterize the pathogenicity of Z. tritici isolates. All wheat-Z. tritici pairwise confrontations were evaluated under controlled conditions, in growth chambers at 18°C/22°C night/day and 16 hours light at 300 μmol.m⁻².s⁻¹. The first true leaf of 16-day-old seedlings were marked with a black felt to delimit a 7.5 cm length that was inoculated with a solution of water containing 10⁶ spores.mL⁻¹ and one drop of Tween®20 per 15 mL. The inoculum was applied with a square-tipped flat paintbrush six times on each leaf, repeated twice. After inoculation the plants were placed in transparent polyethylene bags for 72 hours to initiate infection. At 10 dpi, i.e. before the appearance of symptoms, leaves above the inoculated leaf were cut to homogenize light exposure. Visual estimations of necrotic leaf area and sporulating leaf area were done at 14 dpi, 20 dpi and 26 dpi for matrices 12 and 13, and only once at 21 dpi for matrices 14 and 15. For matrices 12 and 13, these observations were used to calculate, for each plant lineage-isolate combination, an area under disease progress curve (AUDPC) for the percentages of necrotic and sporulating leaf areas. The interactions for matrices 12 and 13 were evaluated on three leaves repeated twice in time (total of six leaves) and for

Matrices 16, 17 and 17b: Phytophthora capsici-Capsicum annuum (pepper)

matrices 14 and 15 on three leaves repeated thrice in time (total of nine leaves).

To build matrix 16, the pathogenicity of six isolates of *Phytophthora capsici*, the causal agent of root and crown rot of chilli and bell peppers, collected in pepper fields in Algeria was measured in ten *Capsicum annuum* cultivars (F_1 hybrids or inbred lines) (Messaouda et al. 2015). Six plants per accession were inoculated by depositing a plug of 4 mm in diameter of mycelium of *P. capsici* cultivated on V8 medium on the fresh section of the primary stem extemporaneously decapitated (Lefebvre and Palloix 1996). Inoculated plants were kept in a growth chamber under controlled conditions with 12h photoperiod, a temperature of $22 \pm 2^{\circ}$ C and 100% relative humidity. *P. capsici* progresses to the

609

610

611

612

613

614

615

616

617

618

619

620

621622

623

624

625

626 627

628

629

630

631

632

633

634

635

636

637

638

639

640

641

bottom of the stem causing a necrosis of the stem. The length of stem necrosis at 15 dpi is reported in matrix 16. For matrix 17, 53 accessions of C. annuum were inoculated by six isolates of P. capsici. The C. annuum accessions originated from 20 countries from America, Europe, Asia and Africa, and included accessions that had different levels of partial resistance to isolate P. capsici 'Pc101' and a few susceptible accessions. The six P. capsici isolates were isolated from pepper plants in France and Turkey, were of A1 mating type and differed in pathogenicity. A minimum of six plants per accession, seven-eight week old, were inoculated as described for matrix 16. Inoculated plants were kept in a growth chamber under controlled conditions with a photoperiod of 12h at 24°C under artificial light and 22°C at obscurity. The length of stem necrosis was measured six times from three to 21 dpi and the Area Under the Disease Progress Curve (AUDPC) of necrosis length was considered in matrix 17. Because matrix 17 contained a large number of zero-values cells, matrix 17b was derived by withdrawing redundant columns and columns entirely made of zero-valued cells. Matrix 18: Phytophthora infestans-Solanum lycopersicum Matrix 18 was built by inoculating eight Solanum sp. accessions with seven isolates of Phytophthora infestans, the causal agent of tomato late blight. The accessions consisted of three inbred lines of cultivated tomato (Solanum lycopersicum) and five accessions of the wild relative species S. pimpinellifolium, S. habrochaites and S. pennellii. Some of them are known to carry the Ph-1, Ph-2 or Ph-3 genes, controlling resistance to races 0, 1 and 2 of P. infestans, respectively. The P. infestans isolates were collected on tomato or potato plants in France and Poland and were chosen because they varied in mating type (A1 or A2) and differed in pathogenicity. Mycelium was grown on pea juicebased agar medium for 10 days and six plants per accession, 3-4 week old, were inoculated using the protocol described for matrix 16 (Danan et al. 2009). Inoculated plants were kept in a growth chamber under controlled conditions with a photoperiod of 14h at 21°C under artificial light and 17°C at obscurity. High humidity was maintained by artificial mist. Stem necrosis length was scored four times from three to 14 dpi and the AUDPC was calculated. Matrix 19: Aphanomyces euteiches-Fabaceae (pea, vetch, faba bean, alfalfa) Eight accessions from four leguminous species (pea, alfalfa, vetch, faba bean), which previously showed various levels of resistance, were inoculated with 34 Aphanomyces euteiches isolates sampled from the main French pea growing regions in a growth chamber (thermo period: 25/23°C and 16h photoperiod). Seven-day-old plants (5 plants * 4 replicates * 2 experiments for each accession-isolate combination) were inoculated by applying 5 mL of a zoospore suspension adjusted

643

644

645

646

647

648

649

650

651

652

653

654

655

656

657

658

659

660

661

662

663

664

665

666

667

668

669670

671

672673

to 5.10³ spores / mL. After inoculation, the vermiculite substrate was saturated with water to provide favorable conditions for infection. After 10 days, the plants were carefully removed from the vermiculite, the roots were washed in tap water and disease severity (DS) was scored on each plant using a 0–5 scale: 0 = no symptoms; 1 = traces of discoloration on the roots (<25%); 2 = discoloration of 25 to 50% of the roots; 3 = discoloration of 50 to 75% of the roots; 4 = discoloration of >75% of the roots; 5 = dead plant. ANOVA was performed with the DS score as the dependent variable, the A. euteiches isolate and the plant accession as fixed factors and the replicate and experiment as random factors. From the ANOVA, least square means (LSmeans) were calculated for each A. euteiches isolate-plant accession combination. In the present study, LSmeans values of root DS scores were analysed. More details are provided in Quillévéré-Hamard et al. (2018). Matrix 20: Aphanomyces euteiches-Pisum sativum Ten pea accessions were inoculated with 43 A. euteiches isolates sampled from the main French pea growing regions in a growth chamber. The ten pea accessions consisted of (i) eight Near-Isogenic-Lines (NILs) carrying one, two, three or five resistance alleles at main QTLs, in a common genetic background and (ii) two control lines, including one susceptible variety and one highly resistant line. The experimental design, inoculation procedure, disease scoring scale and statistical analysis were similar to that described for matrix 19, except for inoculum concentration (2.10² spores / mL) and the scoring date (seven days after inoculation). LSMean values of root DS scores were used. More details are provided in Quillévéré-Hamard et al. (2020). Matrices 21 and 22: Plasmopara viticola-Vitis vinifera (grapevine) A set of 33 Plasmopara viticola strains, the causal oomycete of grapevine downy mildew, was inoculated on eight grapevine varieties. The host panel was constituted of seven grapevine varieties carrying the main resistance factors currently used in European breeding programs (Rpv1, Rpv3.1, Rpv3.2, Rpv5, Rpv6, Rpv10 and Rpv12) and one susceptible variety (Chardonnay). Cuttings from these varieties were grown in a glasshouse under natural photoperiod. Each strain-variety combination was replicated on five leaf discs from five different plants that were excised in the fourth leaf below the apex. Leaf discs were sprayed with 4 mL of a suspension of 10⁵ / mL sporangia of *P. viticola*. They were incubated in a climatic chamber for six days at 18°C with 12h/12h light/dark photoperiod. At six dpi, necrosis was rated on a scale of 0 to 4, based on the number of necroses counted per leaf disk (0 = no necrosis; 1 = <10 necroses; 2 = from 10 to 30 necroses; 3 = from 30 to 60 necroses; 4 = >60

675

676

677

678

679680

681

682

683

684

685

686

687

688

689

690

691

692

693

694

695

696

697

698

699700

701

702

703

704

705

necroses) (matrix 21) and sporulation was assessed on leaf discs by automatic image analysis (number of black pixels on the total leaf disc area) (matrix 22). Matrices 23 and 24: Aphis gossypii-Cucumis melo Matrices 23 and 24 were obtained through assessment of the resistance of 13 melon accessions to nine aphid (Aphis gossypii) clones (Boissot et al. 2016). The host panel consisted in twelve partiallyresistant lines originating from Africa, India, China, Asia and Far East Asia, Mediterranean basin and North America and a susceptible cultivar originating from Mediterranean basin. Two lines were wild accessions and the others from breeding programs. They contained at least one to three homologs of Vat, a gene conferring resistance to A. gossypii. The 13 melon lines belonged to three genetic groups representative of melon diversity (Boissot et al., submitted). The aphid panel consisted in nine clones collected in France and French West Indies. Except clone NM1 that was observed on plant species belonging to six families, the clones have been observed exclusively (or almost exclusively) on cucurbit plants and belong to the same genetic cluster. For phenotyping, ten adult aphids were deposited on melon plantlets. Three days later, the number of aphids remaining on the plantlets was recorded as the 'Acceptance' parameter (matrix 23). Seven days after aphid deposition, the adults were counted, and the density of nymphs was estimated on a scale of 0 to 6. The 'Colonization' parameter was calculated as [density of nymphs + In(number of adults + 0.001)] (matrix 24). The 'Acceptance' and 'Colonization' parameters were collected for at least eight plantlets of each melon accession. Each test was conducted with one aphid clone on a subset of melon accessions. Matrices 25 and 26: Globodera pallida-Solanum tuberosum (potato) Matrix 25 was obtained through the inoculation of 20 populations of the potato cyst nematode Globodera pallida on ten potato accessions. Those potato accessions were characterized by different levels of quantitative resistance. A susceptible potato cultivar, Désirée, was also used as a control. Among the 20 G. pallida populations, 14 came from South-America (Peru and Chile) and six from Europe. To perform G. pallida inoculation, ten cysts were locked in a tulle bag and placed in a pot three-quarter filled with a soil mixture free of cysts (2/3 sand and 1/3 natural field soil). Four replicates were performed for each potato accession - G. pallida population combination, i.e. for each *G. pallida* population, four bags were inoculated to four tubers of the same potato accession. One potato tuber was planted per pot and covered with the same soil mixture. Potato plants grew in

the greenhouse, under controlled conditions (15°C night during 8h and 20°C day during 16h), for 120 days. After 120 days, newly formed cysts were extracted from the soil, using a Kort elutriator. The number of newly formed cysts was counted using a magnifying stereomicroscope, and divided by the number of newly formed cysts produced on the susceptible cultivar Désirée (relative value).

For matrix 26, the measured fitness trait was the hatching of cysts which is induced by host root exudates. It was produced using a cross-hatching assay between 13 populations of *G. pallida* and root exudates from 12 wild potato accessions, belonging to species *Solanum huancabambense*, *S. mochiquense*, *S. sogarandinum*, *S. ambosinum*, *S. medians*, *S. pampasense*, *S. santalallae*, *S. marinasense*, *S. sparsipilum*, *S. raphanifolium*, *S. limbaniense* and *S. leptophyes*, to test the hypothesis of local adaptation between Peruvian *G. pallida* populations and Peruvian wild potato accessions (Gautier et al. 2020). All details about *G. pallida* populations, root-exudates and the *in vitro* hatching assay are available in Gautier et al. (2020). Briefly, three cysts of each population were put on a sieve in 1.5 mL of root exudates (with four to five replicates) and after 30 days, the number of hatched juveniles was counted. At the end of the experiment, cysts were crushed and the number of unhatched viable eggs was counted, in order to calculate a hatching percentage.

Matrices 27 to 32: Potato virus Y-Capsicum annuum

The *Capsicum annuum* accessions were doubled-haploid lines issued from the F₁ hybrid between accessions Perennial, carrying several *Potato virus Y* (PVY) resistance QTLs, and the susceptible accession Yolo Wonder. They were chosen based on the lack of a major-effect resistance gene but contrasted levels of quantitative resistance (Caranta et al. 1997). The PVY populations were issued from cDNA clones of isolates SON41p and LYE84.2 and recombinants between these two cDNA clones (Montarry et al. 2012). *Capsicum annuum* accessions were mechanically inoculated with the different PVY populations and the virus load at the systemic level was estimated one month post inoculation by quantitative DAS-ELISA as described in Quenouille et al. (2014) (matrices 27 and 28). In addition, the area under the disease progress curve (AUDPC) was calculated using a semi-quantitative scoring scale as in Caranta et al. (1997) (matrices 29 and 30) and the dry weight of infected relative to mock-inoculated plants was estimated as in Montarry et al. (2012) (matrices 31 and 32). Matrices 27, 29 and 31 on one side and matrices 28, 30 and 32 on the other side correspond to two independent experiments with slightly different sets of PVY populations.

Network analyses

739

740

741

742

743744

745

746

747

748

749

750

751

752

753

754

755

756

757

758

759

760

761

762

763

764

765

766

767

768

769

770

771

The nestedness and modularity of the different matrices were estimated, and their statistical significance tested respectively with the 'bipartite' and 'igraph' packages of the R software version 3.5.1 (http://cran.r-project.org/). These analyses were initially developed for the study of social, then of ecological, networks (or equivalently matrices) containing counts of links between individuals or between interacting species. Hence, to perform these analyses, the matrices should only contain integer values. Moreover, some nestedness or modularity algorithms cannot run in the absence of zero-valued matrix cells or in the presence of an excess of zero-valued cells leading to an unconnected network. Consequently, the first step consisted in transforming the actual matrices accordingly. In all matrices, pathogenicity trait values were transformed into integers from 0 to 9. For this, ten intervals with equal sizes and spanning the range of the pathogenicity trait values of the actual matrix were defined. The bounds of these intervals are $[P_{min} + (P_{max} - P_{min})^* i/10$, $P_{min} + (P_{max} - P_{min})^* (i + 1)/10]$, with i being an integer in the [0,9] interval and P_{max} and P_{min} being the maximal and minimal pathogenicity trait values in the whole matrix, respectively. Then, depending on its inclusion in a given pathogenicity trait value interval defined as above, each matrix value was transformed into the corresponding i integer value. When necessary, the matrix was modified in order that grades 0 and 9 correspond to the minimal and maximal pathogenicity classes, respectively, and not the opposite. A continuous distribution of the pathogenicity grades was observed in 30 of 32 matrices (Fig. 2). However, for matrices 17b and 22 that contained a large number of zero-values cells, phenotypic values were log-transformed to spread out the data more evenly among the ten phenotypic classes. As these log-transformed matrices produced similar results to the actual matrices in terms of significance of nestedness and modularity, only the latter are shown. To test if the matrices could fit with the inverse-matching-allele model (Fig. 1), we also analyzed the "reverse matrices", where 0 and 9 correspond to the minimal and maximal plant resistance classes, respectively. Methods to estimate nestedness and modularity are detailed in Weitz et al. (2013). Whereas many algorithms can measure the nestedness of matrices containing binary data (0 and 1), only two algorithms were available for matrices containing quantitative numeric data: the weighted nestedness metric based on overlap and decreasing filling (wNODF algorithm) (Almeida-Neto et al. 2008) and the weightedinteraction nestedness estimator (WINE algorithm) (Galeano et al. 2009). In the R software, the 'nested' and 'wine' functions were used to estimate the wNODF and WINE scores, respectively. Because none of the module detection algorithms developed to date provide consistently optimal results in all matrices (Aldecoa and Marín 2013), we used seven different algorithms implemented into the R software (see Supplementary Methods 1 for details). To determine the statistical significance of the patterns (nestedness or modularity) of the plant-parasite interaction matrices, the

773

774

775

776

777

778

779

780

781

782

783

784

785

786

787

788 789

790

791

792

793

794

795

796

797

798

799

800

801

802

803

experiments.

actual interaction matrices were compared to matrices simulated under several null models (Supplementary Methods 1). As the modularity algorithms (and nestedness algorithms to a lower extent) and null models provided contrasted results (Tables 2 to 4), we used simulations to compare their performances (type I and type II error rates) and help the interpretation of the results (Supplementary Methods 1; Tables S1 to S18). Two modularity algorithms (walktrap and label prop) provided modularity estimates of 0 (or near 0) for almost all actual matrices and associated null models. Moreover, almost all simulations also provided modularity estimates of 0 with these algorithms, hampering the evaluation of type I and type II error rates (Supplementary Methods 1). Consequently, these two algorithms were not considered for further analyses. Acknowledgements Marie-Claire Kerlan and Lionel Renault are acknowledged for there help to produce matrix number 25 and Anne Massire, Ghislaine Nemouchi, Thérèse Phaly, Bruno Savio and Patrick Signoret for their assistance to produce matrix number 17. We thank Amine Slim from the National Gene Bank of Tunisia (NGBT) for providing seeds of the durum wheat landrace "Mahmoudi Journine" used to build the matrices 12 and 13, and we thank Aurélie Ducasse and Johann Confais for their help in acquiring phenotypic data on the wheat-Zymoseptoria tritici pathosystem found in matrices 14 and 15. We thank Isabelle Demeaux (INRAE, SAVE) for providing technical assistance with the downy mildew/grapevine pathosystem. Anne Quillévéré-Hamard, Gwenola Le Roy and Christophe Le May are acknowledged for having co-supervised, managed and/or significantly contributed to the production of matrices number 19 and 20. We thank Loup Rimbaud and Emmanuel Szadkowski (INRAE, PACA) for their comments on an earlier version of the manuscript and Michel Pitrat (INRAE, PACA) for his help for analyses of matrices 10 and 11. We thank the staff of the INRAE CRB-Leg (https://www6.paca.inrae.fr/gafl/CRB-Legumes) who maintained the pepper and melon germplasm collections of the GAFL research unit, and of the INRAE experimental facilities of the Plant Pathology research unit (https://doi.org/10.15454/8DGF-QF70), the GAFL experimental unit and the PHENOTIC core facility in Angers (https://doi.org/10.15454/U2BWFJ) who ensured the production of the plants and maintenance of plant-growth facilities that allowed us to do this work. We thank the staff of the INRAE experimental facilities of IGEPP for having provided and managed equipment for the

805

806 807

808

809

810

811

812813

814

815

816

817

818

819

820

821

822

823

824

825

Funding The research was supported by the French National Research Agency (ANR) programs BIOADAPT (grant no. ANR-12-ADAP-0009-04), ArchiV (grant no. ANR-18-CE32-0004-01), CEDRE (grant no. ANR-05-PADD-05) and PeaMUST (grant no. ANR-11-BTBR-0002), the PROGRAILIVE project (grant RBRE160116CR0530019) funded by the Bretagne region, France and European FEADER grants, the fundings of the Institut Carnot PLANT2PRO and the Comité Interprofessionnel des Vins de Bordeaux (CIVB), the INRAE departments "Santé des Plantes et Environnement" (project APÔGÉ and PhD thesis of Safa Ben Krima) and "Génétique et Amélioration des Plantes", the INRAE métaprogramme SMaCH (Sustainable Management of Crop Health), the French Ministry of Agriculture and Food for projects "Recherche et mise au point de méthodes pour évaluer des résistances variétales durables à des agents pathogènes" (CTPS project C2008-29), "Nouvelles sources de résistance à Aphis gossypii chez le melon" (CTPS project C06/02) and "Caractérisation de la virulence de Podosphaera xanthii, agent causal de l'oïdium du melon, et développement d'un système de codification des races" (CTPS project C-2012-10). UMR1290 BIOGER benefits from the support of Saclay Plant Sciences-SPS (ANR-17-EUR-0007). **Conflict of interest disclosure** The authors of this manuscript declare that they have no financial conflict of interest with the content of this article. Benoît Moury in one of the Peer Community In Evolutionary Biology recommenders.

Table 1: Datasets used to analyze the structure of quantitative plant-parasite interaction matrices. AUDPC: Area under the disease progress curve.

Matrix number	Parasite	Host plant	Matrix size (host × parasite)	Phenotype	Multiple R ^{2a}	Reference or source
Bacterium						
1	Pseudomonas syringae	Prunus armeniaca (apricot)	20 × 9	Length of flat zone around the inoculation point at the surface of the shoot	0.69	Omrani <i>et al.,</i> unpublished
2	P. syringae	P. armeniaca	20 × 9	Length of flat zone around the inoculation point at the surface of the shoot	0.84	Omrani et al., unpublished
3	P. syringae	P. armeniaca	20 × 9	Length of browning zone around the inoculation point below the bark of the shoot	0.68	Omrani et al., unpublished
4	P. syringae	P. armeniaca	20 × 9	Length of browning zone around the inoculation point below the bark of the shoot	0.75	Omrani et al., unpublished
Fungi						
5	Puccinia hordei	Hordeum vulgare (barley)	12 × 14	Relative latent period	0.84	González et al., 2012
6	Venturia inaequalis	Malus domestica (apple)	14 × 10	% sporulating leaf area (AUDPC)	0.64	Laloi <i>et al.,</i> 2017
7	V. inaequalis	M. domestica	12 × 14	% sporulating leaf area (AUDPC)	0.49	Caffier <i>et al.</i> , 2016
8	V. inaequalis	M. domestica	8 × 24	% sporulating leaf area	0.71	Caffier <i>et al.</i> , 2014
9	Botrytis cinerea	Tomato ^b	12 × 94	Lesion size on leaves	0.59	Soltis <i>et al.</i> , 2019
10	Podosphaera xanthii	Cucumis melo (melon)	19 × 26	Sporulation surface on leaf disks	0.93	Dogimont et al., unpublished
11	P. xanthii	C. melo	19 × 31	Sporulation surface on plants	0.94	Dogimont et al., unpublished
12	Zymoseptoria tritici	Triticum turgidum subsp. durum (durum wheat)	12 × 15	% necrotic leaf area (AUDPC)	0.77	Marcel et al., unpublished
13	Z. tritici	T. turgidum subsp. durum	12 × 15	% sporulating leaf area (AUDPC)	0.76	Marcel et al., unpublished
14	Z. tritici	<i>Triticum aestivum</i> (bread wheat)	15 × 98	% necrotic leaf area	0.63	Marcel et al., unpublished
15	Z. tritici	T. aestivum	15 × 98	% sporulating leaf area	0.63	Marcel et al., unpublished

Oomycetes						
16	Phytophthora capsici	Capsicum annuum (pepper)	10 × 6	Necrosis length on stem (15 days post inoculation)	0.98	Messaouda et al., 2015
17	P. capsici	C. annuum	53 × 6	Necrosis length on stem (AUDPC)	0.78	Cantet et al., unpublished
17b ^c	P. capsici	C. annuum	42 × 6	Necrosis length on stem (AUDPC)	-	Cantet et al., unpublished
18	Phytophthora infestans	Solanum lycopersicum (tomato), S. pimpinellifolium, S.	8 × 7	Necrosis length on stem (AUDPC)	0.90	Ruellan <i>et al.,</i> unpublished
4.0		habrochaites and S. pennellii	0 05	B	0.05	0 111/ / / 11
19	Aphanomyces euteiches	Fabaceae ^d	8 × 35	Root disease severity	0.85	Quillévéré-Hamard et al., 2018
20	A. euteiches	Pisum sativum	10 × 43	Root disease severity	0.86	Quillévéré-Hamard et al., 2020
21	Plasmopara viticola	Vitis vinifera (grapevine)	8 × 33	Necrosis on leaves	0.76	Paineau and Delmotte, unpublished
22	P. viticola	V. vinifera	8 × 33	Sporulation on leaves	0.80	Paineau and Delmotte, unpublished
Insect						
23	Aphis gossypii	C. melo	13 × 9	Acceptance of plants	0.61	Boissot et al., 2016
24	A. gossypii	C. melo	13 × 9	Ability to colonize plants	0.59	Boissot et al., 2016
Nematode						
25	Globodera pallida	Solanum tuberosum (potato)	10 × 20	Cyst number (relative values)	0.58	Fournet et al., unpublished
26	G. pallida	Wild potato species	12 × 13	Cyst eclosion rate	0.81	Gautier et al., 2020
Virus						
27	Potato virus Y (PVY)	C. annuum	7 × 8	Virus load	0.79	Doumayrou et al., unpublished
28	PVY	C. annuum	9 × 7	Virus load	0.73	Doumayrou et al., unpublished
29	PVY	C. annuum	8 × 7	Symptom intensity (AUDPC)	0.78	Doumayrou et al., unpublished
30	PVY	C. annuum	7 × 7	Symptom intensity (AUDPC)	0.81	Doumayrou et al., unpublished
31	PVY	C. annuum	8 × 7	Relative dry matter weight	0.66	Doumayrou et al., unpublished
32	PVY	C. annuum	7 × 7	Relative dry matter weight	0.45	Doumayrou et al., unpublished

^a Fit of the linear model: pathogenicity ~ 'parasite strain' + 'plant accession' (multiple coefficient of determination).

^b Two species: cultivated tomato (*Solanum lycopersicum*) and wild tomato (*S. pimpinellifolium*).

^c Matrix 17b is identical to matrix 17 except that columns entirely made of zero-valued cells and redundant columns were removed.

^d Four species: three pea (*Pisum sativum*) accessions, two vetch (*Vicia sativa*) accessions, two faba bean (*Vicia faba*) accessions and one alfalfa (*Medicago sativa*) accession.

Table 2: Analysis of nestnedness of plant-parasite interaction matrices with two methods.

WINE method								wNODF method								
Matrix	Nestedness				Null model	b			Nestedness Null model ^b							
number	$score^a$	В	N	C1	R1	S	C2	R2	score	В	N	C1	R1	S	C2	R2
1	0.78	0 ^c	0	0	0	0	0	0	30.9	0.05	0	0	0	0	0	0
2	0.81	0.01	0	0	0	0	0	0	43.7	0.06	0	0	0	0	0	0
3	0.82	0	0	0	0	0	0	0	47.6	0	0	0	0	0	0	0
4	0.82	0	0	0	0	0	0	0	46.9	0	0	0	0	0	0	0
5	0.83	0	0	0	0	0	0	0	23.2	0.24	0	0.42	0	0	0.36	0
6	0.70	0.12	0	0	0	0	0	0	55.4	0.15	0	0	0	0	0	0
7	0.60	0.49	0	0	0	0	0	0	43.9	0.41	0	0	0	0	0	0
8	0.58	0.01	0	0	0	0	0	0	49.4	0.66	0	0	0	0	0	0
9	0.46	0	0	0	0	0	0	0	42.4	0.01	0	0	0	0	0	0
10	1.01	0	0	0	0	0	0	0	72.7	0	0	0	0	0	0	0
11	1.04	0	0	0	0	0	0	0	75.4	0	0	0	0	0	0	0
12	0.69	0	0	0	0	0	0	0	10.1	0.68	0.85	0.67	1	0.73	0.54	0.90
13	0.73	0	0	0	0	0	0	0	30.0	0.16	0.38	0.48	0.07	0.35	0.39	0.52
14	0.76	0	0	0	0	0	0	0	22.3	0.76	1	1	1	0.06	1	0.22
15	0.68	0.12	0	0	0	0	0	0	41.9	0.31	0	0	0	0	1	0
16	0.84	0	0	0	0	0	0	0	38.6	0.09	0	0	0	0	0	0
17	0.84	0	0	0	0	0	0	0	51.0	0	0	0	0	0	0	0
18	0.93	0	0	0	0	0	0	0	59.2	0	0	0	0	0	0	0
19	0.79	0	0	0	0	0	0	0	9.4	0.89	0.01	0.01	0.80	0.04	0	0.01
20	0.80	0	0	0	0	0	0	0	15.0	0.98	0	0	0	0	0.12	0
21	0.75	0.90	0	0.07	0	0	0.46	0	27.6	0.33	0.42	0.68	0	0.51	1	0.59
22	0.91	0	0	0	0	0	0	0	49.2	0	0	0	0	0	0	0
23	0.75	0	0	0	0	0	0	0.04	6.1	0.65	0.82	0.28	1	0.82	0.48	0.85
24	0.68	0.12	0	0	0.01	0	0	0.02	22.4	0.29	0.98	0.96	0.72	0.95	0.93	0.72
25	0.82	0.01	0	0	0	0	0	0	44.1	0.20	0	0	0	0	0.02	0
26	0.86	0	0	0	0	0	0	0	34.3	0	0	0	0	0	0	0
27	0.69	0.37	0	0	0	0	0	0	51.2	0.41	0	0	0.01	0	0.01	0
28	0.63	0.46	0	0.01	0	0	0	0	58.0	0.27	0	0	0	0	0	0
29	0.78	0.01	0	0	0	0	0	0	22.3	0.97	0.81	0.62	0.02	0.88	0.96	0.18
30	0.84	0.03	0	0	0	0	0	0	66.8	0.01	0	0	0	0	0	0
31	0.77	0	0	0	0	0	0	0.01	21.3	0.05	0.30	0.49	0.04	0.21	0.35	0.06
32	0.52	0.31	0.06	0	0.36	0.06	0.01	0.71	11.1	0.13	0.65	0.55	0.90	0.62	0.51	0.74

^aMean of 100 estimates.

^bSee Supplementary Methods 1 for details of the null models.

cNestedness significance: the probability value (p-value) indicates the frequency of null-model matrices showing a strictly higher nestedness score than that of the actual matrix. P-values ≤ 0.05 (significant nestedness) are in bold on grey cells and p-values > 0.95 (significant anti-nestedness) are in white on black cells.

Table 3. Analysis of modularity of plant-parasite interaction matrices with the *spinglass* method.

				Spingla	SS				
Matrix		Number of				Null model ^c			
number	Modularity score ^a	$modules^b$	В	N	C1	R1	S	C2	R2
1	0.058	-	0.69 ^d	0.76	0.79	0.71	0.93	0.89	0.93
2	0.070	-	0.86	1	1	0.84	1	1	0.92
3	0.084	-	0.32	0.68	0.38	0.42	0.36	0.20	0.2
4	0.095	-	0.95	1	1	0.97	0.99	0.98	0.8
5	0.070	2	0	0	0	0	0	0	0
6	0.102	3	0	0	0	0	0	0	0
7	0.079	-	0.97	1	1	1	0.99	0.96	0.9
8	0.069	-	0.07	0.53	0.35	0.05	0.70	0.66	0.3
9^e	0.069	-	0.96	1	1	0.97	0.89	0.97	0.8
10	0.077	3	0	0.07	0	0.03	0.01	0.05	0.0
11	0.086	3	0	0	0	0	0	0	0
12	0.057	-	0.98	0.98	0.99	0.96	1	0.99	0.9
13	0.072	-	0.08	0.27	0.23	0.06	0.73	0.43	0.3
14	0.062	2	0	0	0	0	0	0.01	0
15 ^e	0.079	-	0.06	0.61	0.14	0.68	0.13	0.08	0.3
16	0.078	-	0.20	0.39	0.32	0.37	0.66	0.41	0.8
17b ^f	0.097	3	0.05	0.25	0.15	0.06	0.05	0.01	0.0
18	0.097	-	0.77	0.98	0.97	0.93	0.95	0.71	0.9
19	0.051	-	1	1	1	1	1	1	1
20	0.040	-	1	1	1	1	1	1	1
21	0.092	-	0.44	0.97	0.59	1	0.99	0.56	0.9
22	0.083	-	0.15	NA^g	0.06	0.44	NA^g	0.02	0.3
23 ^e	0.072	-	0.41	0.39	0.27	0.36	0.65	0.57	0.5
24	0.063	-	0.30	0.52	0.58	0.27	0.81	0.77	0.5
25	0.091	-	0.75	0.99	0.78	0.99	0.75	0.58	0.4
26 ^e	0.045	-	0.79	0.91	0.86	0.90	1	0.99	1
27	0.130	-	0.73	0.83	0.74	0.85	0.76	0.74	0.8
28	0.090	-	0.96	0.98	0.98	1	0.91	1	0.9
29 ^e	0.078	-	0.89	0.79	0.91	0.79	0.93	1	0.8
30	0.095	-	0.49	0.80	0.82	0.56	0.59	0.58	0.4
31	0.061	-	0.98	1	0.98	1	1	1	0.9
32	0.065	-	1	0.99	1	0.99	0.97	0.97	0.9

^aMaximum of 100 estimates.

^bThe optimal number of modules determined by *spinglass* is indicated only for matrices significantly modular with a majority of null models (Fig. 4).

^cSee Supplementary Methods 1 for details of the null models.

 d Modularity significance: the probability value (p-value) indicates the frequency of null-model matrices showing a strictly higher modularity score than that of the actual matrix. P-values ≤ 0.05 (significant modularity) are in bold on grey cells. Significant anti-modularity, when ≤ 5% of null-model matrices show a strictly lower modularity degree than that of the actual matrix, are indicated in white on black cells. Note that some of the indicated p-values are ≥ 0.95 but do not correspond to significant anti-modularity because the modularity degrees of the actual matrix and of some null-model matrices are identical.

eRows and/or columns entirely made of zero-valued cells were removed since the spinglass method cannot estimate the modularity under such circumstances (unconnected graphs).

^fMatrix 17b is identical to matrix 17 except that columns entirely made of zero-valued cells and redundant columns were removed.

^gNA: not available; many null-model matrices had rows and/or columns entirely made of zero-valued cells and the *spinglass* method could not estimate their modularity.

Table 4. Analysis of modularity of plant-parasite interaction matrices with four methods. Only the three null models S, C2 and R2 that provided the lowest rates of false positive modularity in our performance study (Supplementary Methods 1) are presented.

	Ed		Fast gree	edy			Louvai	n		Leading eigenvector						
Matrix	Modularity	N	Iull model ^t)	Modularity	1	Null mode	l b	Modularity	Null model ^b			Modularity	Null model ^b		b
number	score	S	C2	R2	score	S	C2	R2	score	S	C2	R2	score ^a	S	C2	R2
1	0.012	1 ^c	1	0.99	0.074	1	0.99	0.51	0.073	1	1	1	0.061	0.87	0.77	0.62
2	0.022	1	1	0.97	0.091	1	1	0.98	0.091	1	1	1	0.088	1	1	0.37
3	0.099	0.88	0.53	0.06	0.126	1	0.96	0.38	0.127	1	1	0.64	0.116	0.85	0.62	0.12
4	0	1	1	1	0.113	1	1	1	0.115	1	1	1	0.113	1	1	0.40
5	0.018	1	0.01	1	0.065	0.99	0	0.99	0.069	1	0	1	0.069	0.58	0	0.57
6	0.085	1	0.99	1	0.209	0.88	0.03	0.17	0.209	0.96	0.10	0.42	0.206	0.50	0.01	0.07
7	0.138	0.96	0.45	0.79	0.240	1	0.84	1	0.240	1	0.98	1	0.160	1	0.97	0.99
8	0.003	1	1	1	0.121	1	1	0.98	0.125	1	1	1	0.116	0.99	0.97	0.43
9	0	1	1	1	0.074	1	1	1	0.069	1	1	1	0.061	0.99	0.84	0.59
10	0	1	1	1	0.073	1	1	1	0.073	1	0.98	1	0.072	1	0.40	0.47
11	0	1	1	1	0.085	1	0.98	1	0.091	1	0.93	1	0.058	1	0.96	0.99
12	0.004	1	1	0.50	0.041	1	1	0.16	0.044	1	1	0.98	NA^c	NA ^d	NA^d	NA^d
13	0.004	1	1	0.95	0.061	1	1	0.56	0.063	1	1	0.88	0.039	0.96	0.94	0.64
14	0	1	1	1	0.044	0.87	0.04	0.35	0.046	0.48	0.04	0.15	0.010	1	0.41	1
15	0	1	1	1	0.089	1	0.98	1	0.087	1	1	1	0.063	0.99	0.64	1
16	0.007	1	1	1	0.032	1	1	1	0.036	1	1	1	0.022	0.99	0.92	1
17	0.071	1	0.99	1	0.172	1	1	1	0.177	1	1	1	0.155	1	0.99	0.99
17b ^e	0.040	1	1	0.99	0.155	1	1	1	0.158	1	1	1	0.131	0.38	0.21	0.96
18	0.079	1	_ 1	1	0.186	1	1	0.35	0.186	0.99	1	0.16	0.169	0.98	1	0.32
19	0	1	1	1	0.028	1	1	1	0.031	1	1	1	0	1	1	1
20	0	1	1	1	0.029	1	1	. 1	0.029	1	1	1	0.029	0.97	0.88	0.83
21	0.036	1	0.23	1	0.110	1	0	1	0.109	1	0.01	1	0.096	0.99	0	1
22	0.020	1	0.98	1	0.165	1	0.89	1	0.165	1	0.93	1	0.157	1	0.27	1
23	0.011	1	1	0.88	0.068	0.91	0.81	0.22	0.070	0.99	0.99	0.70	0.058	0.67	0.44	0.26
24	0.035	0.99	0.86	0.91	0.092	0.79	0.75	0.20	0.095	0.94	0.93	0.60	0.066	0.90	0.82	0.74
25	0.066	1	0.35	1	0.167	1	0.49	1	0.168	1	0.84	1	0.118	1	0.78	1
26	0	1	1	1	0.041	1	1	1	0.046	1	1	1	0.044	0.82	0.39	0.84
27	0.034	0.96	0.86	0.69	0.102	1	1	1	0.103	1	1	1	0.086	1	1	1
28	0	1	1	1	0.151	1	1	1	0.153	1	1	1	0.108	1	1	1
29	0.008	1	1	1	0.069	1	1	1	0.069	1	1	1	0.048	1	0.95	1
30	0.043	0.98	0.91	0.91	0.087	1	1	1	0.092	1	1	1	0	1	1	1
31	0.013	0.98	0.97	0.65	0.062	1	1	0.89	0.062	1	1	1	0.055	0.69	0.71	0.59
32	0.058	0.70	0.84	0.72	0.062	0.95	0.99	0.59	0.062	1	1	1	0.022	0.97	0.96	0.73

^aMaximum of 100 estimates.

^bSee Supplementary Methods 1 for details of the null models.

 $^{\circ}$ Modularity significance: the probability value (p-value) indicates the frequency of null-model matrices showing a strictly higher modularity score than that of the actual matrix. P-values ≤ 0.05 (significant modularity) are in bold on grey cells. Significant anti-modularity, when ≤ 5% of null-model matrices show a strictly lower modularity degree than that of the actual matrix, are indicated in white on black cells. Note that some of the indicated p-values are ≥ 0.95 but do not correspond to significant anti-modularity because the modularity degrees of the actual matrix and of some null-model matrices are identical.

^dNA: not available; the *leading eigenvector* algorithm failed to converge.

^eMatrix 17b is identical to matrix 17 except that columns entirely made of zero-valued cells and redundant columns were removed.

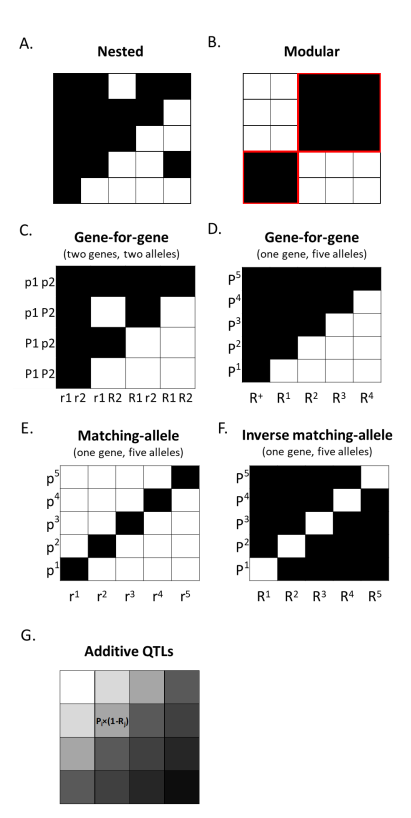


Figure 1. Matrices corresponding to different mechanistic, genetic and evolutionary models of qualitative or quantitative host-parasite interactions. In each case, host genotypes correspond to different columns and parasite genotypes to different rows) and black and white cells (or 1 and 0 grades) correspond to infection or lack of infection, respectively. A: Illustration of an imperfectly nested pattern. B: Illustration of a perfectly modular pattern (modules are delimitated with red lines). C and D: Gene-for-gene (GFG) models with partial or perfectly nested patterns. C: Case of two genes with two alleles in both hosts and parasites. Infection occurs only when no elicitor in the

parasite is recognized by a product of the resistance alleles in the host. In the other situations, resistance is induced and there is no infection. D: Case of a single gene with five alleles in both hosts and parasites. Resistance alleles have various levels of specificity: in some plant accessions resistance can be induced by several parasite strains. E: Matching-allele model. Infection occurs only if the product of the pathogenicity allele is recognized by the product of the susceptibility allele in the host. F: Variation of D with higher specificity: resistance is induced by a specific product present in a single parasite genotype. This model was named "inverse matching-allele" model (Thrall et al. 2016) and has an anti-modular structural pattern. G: Additive QTL model with no plant-parasite QTL \times QTL interaction. For each parasite strain i with pathogenicity level P_i and each plant accession j with resistance level R_{ij} , infection score corresponds to $P_i \times (1-R_j)$.

Superscript figures correspond to alleles of a given gene whereas normal font figures correspond to different genes. Matching genes or alleles at resistance and pathogenicity loci in host and parasite genotypes share the same figure. For simplicity, hosts and parasites are considered haploid. R: resistance allele; r or R⁺: susceptibility allele; P: allele controlling lack of pathogenicity; p: pathogenicity allele.

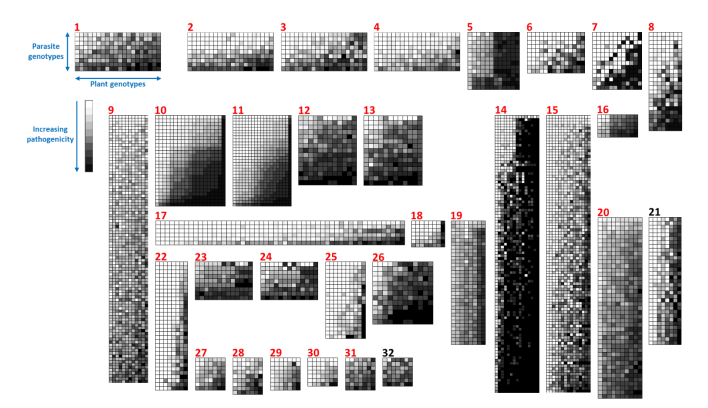


Figure 2. Overview of the 32 analyzed plant-parasite matrices (Table 1). Different plant accessions and parasite strains correspond to different columns and rows, respectively. White to black shades in each cell correspond to an increasing gradient of pathogenicity or infectivity (corresponding to 0 to 9 values in the analysed matrices) for a given plant and parasite pair. Rows and columns were ordered by increasing marginal totals, revealing the nested patterns. Red numbers correspond to significant nestedness (*WINE* algorithm) (Table 2).

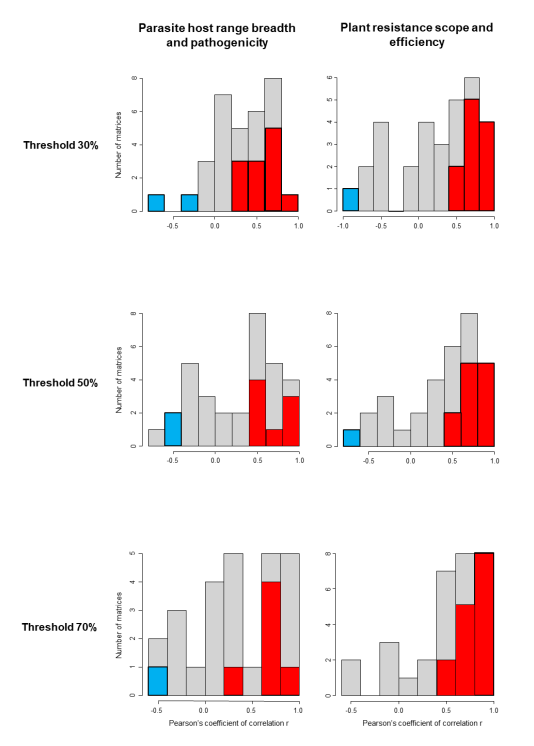


Figure 3. Distributions of Pearson's coefficients of correlation (r) between parasites host range breadth and pathogenicity (left) or between plant resistance efficiency and scope (right) across the 32 analysed matrices for different thresholds separating hosts and non-hosts (or parasites included or not included in the resistance scope). Each threshold corresponds to a percentage of the maximal pathogenicity value in each matrix (only results obtained with thresholds corresponding to 30%, 50% and 70% of the maximal pathogenicity value are shown; results were similar for other thresholds). In blue and red: significantly negative or positive r values (p-value < 0.05). For some thresholds and some matrices, the coefficient of correlation could not be calculated because too few pathogenicity data remained.

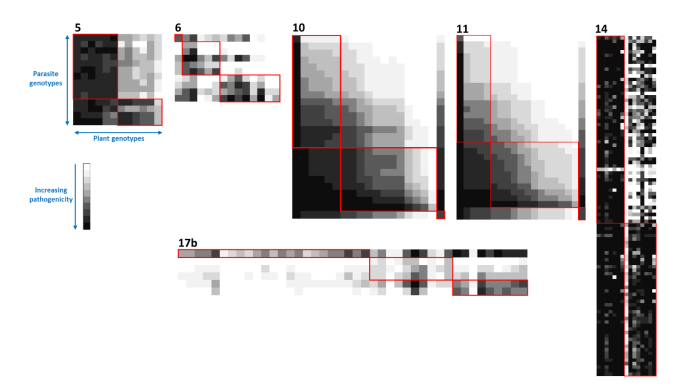


Figure 4. Overview of the six plant-parasite matrices showing significant modularity with the *spinglass* algorithm (Table 3). Rows and columns were ordered by modules, delimited by red lines. See legend of Figure 2 for the representation of matrices.

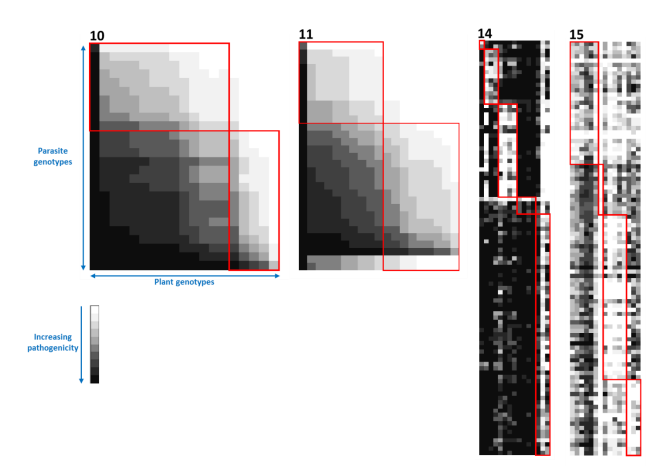


Figure 5. Overview of the four plant-parasite matrices showing significant modularity with the *spinglass* algorithm when matrices were transformed such that 0 values correspond to the maximal plant susceptibility and 9 values to the maximal plant resistance (but note that the matrices are represented such that 0 to 9 values correspond to a plant resistance to susceptibility gradient, as in the original matrices). Rows and columns were ordered by modules, delimited by red lines. See legend of Figure 2 for the representation of matrices.

References

- Aldecoa, R., Marín, I. 2013. Exploring the limits of community detection strategies in complex networks. Sci. Rep. 3:2216.
- Almeida-Neto, M., Ulrich, W. 2011. A straightforward computational approach for measuring nestedness using quantitative matrices. Environ. Model. Softw. 26:173-176.
- Almeida-Neto, M., Guimaraes, P., Guimaraes, P.R., Loyola, R.D., Ulrich, W. 2008. A consistent metric for nestedness analysis in ecological systems: reconciling concept and measurement. Oikos 117:1227–1239.
- Bahri, B., Kaltz, O., Leconte, M., de Vallavieille-Pope, C., Enjalbert, J. 2009. Tracking costs of virulence in natural populations of the wheat pathogen, *Puccinia striiformis* f.sp.tritici. BMC Evol. Biol. 9:26.
- Bartoli, C., Roux, F. 2017. Genome-wide association studies in plant pathosystems: Toward an ecological genomics approach. Front. Plant Sci. 8:763.
- Ben Krima, S., Slim, A., Gelisse, S., Kouki, H., Nadaud, I., Sourdille, P., Yahyaoui, A., Ben M'barek, S., Suffert, F., Marcel, T.C. 2020. Life story of Tunisian durum wheat landraces revealed by their genetic and phenotypic diversity. bioRxiv 2020.08.14.251157.
- Barrett, L.G., Encinas-Viso, F., Burdon, J.J., Thrall, P.H. 2015. Specialization for resistance in wild host-pathogen interaction networks. Front. Plant Sci. 6:761.
- Bascompte, J., Jordano, P., Melián, C.J., Olesen, J.M. 2003. The nested assembly of plant–animal mutualistic networks. Proc. Natl. Acad. Sci. USA 100:9383–9387.
- Bent, A. F., Mackey D. 2007. Elicitors, effectors, and R genes: the new paradigm and a lifetime supply of questions. Annu. Rev. Phytopathol. 45:399-436.
- Blondel, V.D., Guillaume, J.-L., Lambiotte, R., Lefebvre, E. 2008. Fast unfolding of communities in large networks. J. Stat. Mech.: Theory Exp. 10:P10008.
- Boissot, N., Thomas, S., Chovelon, V., Lecoq, H. 2016. NBS-LRR-mediated resistance triggered by aphids: viruses do not adapt; aphids adapt via different mechanisms. BMC Plant Biol. 16:25.
- Brown J.K.M. 2015. Durable resistance of crops to disease: a Darwinian perspective. Annu. Rev. Phytopathol. 53:513-539.
- Brualdi, R.A., Sanderson, J.G. 1999. Nested species subsets, gaps, and discrepancy. Oecologia 119:256-264.
- Bruns, E., Carson, M., May, G. 2014. The Jack of all trades is master of none: a pathogen's ability to infect a greater number of host genotypes comes at a cost of delayed reproduction. Evolution 68:2453-2466.
- Caffier, V., Didelot, F., Pumo, B., Causeur, D., Durel, C.E., Parisi, L. 2010. Aggressiveness of eight *Venturia inaequalis* isolates virulent or avirulent to the major resistance gene *Rvi6* on a non-*Rvi6* apple cultivar. Plant Pathol. 59:1072-1080.
- Caffier, V., Lasserre-Zuber, P., Giraud, M., Lascostes, M., Stievenard, R., Lemarquand, A., Van De Weg, E., Expert, P., Denancé, C., Didelot, F., Le Cam, B., Durel, C.E. 2014. Erosion of quantitative host resistance in the apple Venturia inaequalis pathosystem. Infect. Genet. Evol. 27:481-489.
- Caffier, V., Le Cam, B., Al Rifaï, M., Bellanger, M.N., Comby, M., Denancé, C., Didelot, F., Expert, P., Kerdraon, T., Lemarquand, A., Ravon, E., Durel, C.E. 2016. Slow erosion of a quantitative apple

- resistance to *Venturia inaequalis* based on an isolate-specific Quantitative Trait Locus. Infect. Genet. Evol. 44:541-548.
- Caranta, C., Lefebvre, V., Palloix, A. 1997. Polygenic resistance of pepper to potyviruses consists of a combination of isolate-specific and broad-spectrum quantitative trait loci. Mol. Plant-Microbe Interact. 10:872–878.
- Castagnone-Sereno, P., Bongiovanni, M., Wajnberg E. 2007. Selection and parasite evolution: A reproductive fitness cost associated with virulence in the parthenogenetic nematode *Meloidogyne incognita*. Evol. Ecol. 21:259-270.
- Clauset, A., Newman, M.E.J., Moore, C. 2004. Finding community structure in very large networks. Phys. Rev. E 70:066111.
- Danan, S., Chauvin, J.-E., Caromel, B., Moal, J.-D., Pellé, R., Lefebvre, V. 2009. Major-effect QTLs for stem and foliage resistance to late blight in the wild potato relatives *Solanum sparsipilum* and *S. spegazzinii* are mapped to chromosome X. Theor. Appl. Genet. 119:705–719.
- Delmas, C.E.L., Fabre, F., Jolivet, J., Mazet, I.D., Cervera, S.R., Delière, L., Delmotte F. 2016. Adaptation of a plant pathogen to partial host resistance: selection for greater aggressiveness in grapevine downy mildew. Evol. Appl. 9:709–725.
- Desbiez, C., Gal-On, A., Girard, M., Wipf-Scheibel, C., Lecoq, H. 2003. Increase in *Zucchini yellow mosaic virus* symptom severity in tolerant zucchini cultivars is related to a point mutation in P3 protein and is associated with a loss of relative fitness on susceptible plants. Phytopathology 93:1478-1484.
- Dormann, C.F., Fründ, J., Schaefer, H.M. 2017. Identifying causes of patterns in ecological networks: Opportunities and limitations. Annu. Rev. Ecol. Evol. Syst. 48:559-584.
- Feng, W, Takemoto, K. 2014. Heterogeneity in ecological mutualistic networks dominantly determines community stability. Sci. Rep. 4:5912.
- Fenton, A., Antonovics, J., Brockhurst, M.A. 2009. Inverse-gene-for-gene infection genetics and coevolutionary dynamics. Am. Nat. 174:E230-E242.
- Flor, H.H. 1956. The complementary genic systems in flax and flax rust. Adv. Genet. 8:29-54.
- Flores, C.O., Meyer, J.R., Valverde, S., Farr, L., Weitz, J.S. 2011. Statistical structure of host-phage interactions. Proc. Natl. Acad. Sci. USA 108: E288–97.
- Flores, C.O., Valverde, S., Weitz, J.S. 2013. Multi-scale structure and geographic drivers of cross-infection within marine bacteria and phages. ISME J. 7:520-532.
- Fortuna, M.A., Barbour, M.A., Zaman, L., Hall, A.R., Buckling, A., Bascompte, J. 2019. Coevolutionary dynamics shape the structure of bacteria—phage infection networks. Evolution 73:1001–1011.
- Fournet, S., Eoche-Bosy, D., Renault, L., Hamelin, F.M., Montarry, J. 2016. Adaptation to resistant hosts increases fitness on susceptible hosts in the plant parasitic nematode *Globodera pallida*. Ecol. Evol. 6:2559-2568.
- Fraile, A., Pagán, I., Anastasio, G., Saez, E., García-Arenal, F. 2011. Rapid genetic diversification and high fitness penalties associated with pathogenicity evolution in a plant virus. Mol. Biol. Evol. 28:1425-1437.
- Galeano, J., Pastor, J.M., Iriondo, J.M. 2009. Weighted-Interaction Nestedness Estimator (WINE): A new estimator to calculate over frequency matrices. Env. Model. Soft. 24:1342-1346.

- Gallois, J.-L., Moury, B., German-Retana, S. 2018. Role of the genetic background in resistance to plant viruses. Int. J. Mol. Sci. 19:2856.
- Gautier, C., Fournet, S., Piriou, C., Renault, L., Yvin, J.C., Nguema-Ona, E., Grenier, E., Montarry, J. 2020. Plant-parasite coevolution: a weak signature of local adaptation between Peruvian *Globodera pallida* populations and wild potatoes. Ecol. Evol. 10:4156-4163.
- González, A.M., Marcel, T.C., Niks, R.E. 2012. Evidence for a minor gene-for-minor gene interaction explaining nonhypersensitive polygenic partial disease resistance. Phytopathology 102:1086-1093.
- Gotelli, N.J., Graves, G.R. 1996. Null models in ecology. Smithsonian Institution Press, Washington D.C.
- Grosberg, R.K., Hart, M.W. 2000. Mate selection and the evolution of highly polymorphic self/nonself recognition genes. Science 289:2111-2114.
- Huang, Y.J., Balesdent, M.-H., Li, Z.Q., Evans, N., Rouxel, T., Fitt, B.D.L. 2010. Fitness cost of virulence differs between the *AvrLm1* and *AvrLm4* loci in *Leptosphaeria maculans* (phoma stem canker of oilseed rape). Eur. J. Plant Pathol. 126:279-291.
- Ishibashi, K., Mawatari, N., Miyashita, S., Kishino, H., Meshi, T., Ishikawa, M. 2012. Coevolution and hierarchical interactions of *Tomato mosaic virus* and the resistance gene *Tm-1*. PLoS Pathog. 8:e1002975.
- Janzac, B., Montarry, J., Palloix, A., Navaud, O., Moury, B. 2010. A point mutation in the polymerase of *Potato virus Y* confers virulence towards the *Pvr4* resistance of pepper and a high competitiveness cost in susceptible cultivar. Mol. Plant-Microbe Interact. 23:823-830.
- Jenner, C.E., Wang, X.W., Ponz, F., Walsh, J.A. 2002. A fitness cost for *Turnip mosaic virus* to overcome host resistance. Virus Res. 86:1-6.
- Jonhson, S., Domínguez-García, V., Muñoz, M.A. 2013. Factors determining nestedness in complex networks. PLOS ONE 8:e74025.
- Joppa, L.N., Montoya, J.M., Solé, R., Sanderson, J., Pimm, S.L. 2010. On nestedness in ecological networks. Ecol. Evol. Res. 12:35–46.
- Khatabi, B., Wen, R.H., Hajimorad, M.R. 2013. Fitness penalty in susceptible host is associated with virulence of *Soybean mosaic virus* on *Rsv1*-genotype soybean: a consequence of perturbation of HC-Pro and not P3. Mol. Plant Pathol. 14:885-897.
- Kidner, J., Moritz, R.A.F. 2013. The Red Queen process does not select for high recombination rates in haplodiploid hosts. Evol. Biol. 40:377–84.
- Laloi, G., Vergne, E., Durel, C.-E., Le Cam, B., Caffier, V. 2017. Efficiency of pyramiding of three quantitative resistance loci to apple scab. Plant Pathol. 66:412-422.
- Lambrechts, L. 2010. Dissecting the genetic architecture of host–pathogen specificity. PLoS Pathog. 6:e1001019.
- Leach, J.E., Vera Cruz, C.M., Bai, J.F., Leung, H. 2001. Pathogen fitness penalty as a predictor of durability of disease resistance genes. Annu. Rev. Phytopathol. 39:187-224.
- Lefebvre, V., Palloix, A. 1996. Both epistatic and additive effects of QTLs are involved in polygenic induced resistance to disease: a case study the interaction pepper-*Phytophthora capsici* Leonian. Theor. Appl. Genet. 93:503-511.

- Marcel, T.C., Gorguet, B., Truong Ta, M., Kohutova, Z., Vels, A., Niks, R.E. 2008. Isolate specificity of quantitative trait loci for partial resistance of barley to *Puccinia hordei* confirmed in mapping populations and near-isogenic lines. New Phytol. 177:743-755.
- Messaouda, B., Guechi, A., Mézaache-Aichour, S. 2015. Susceptibility of Algerian pepper cultivars (*Capsicum annuum* L.) to *Phytophthora capsici* strains from different geographic areas. Afr. J. Biotechnol. 14:3011-3018.
- Montarry, J., Hamelin, F.M., Glais, I., Corbière, R., Andrivon, D. 2010. Fitness costs associated with unnecessary virulence factors and life history traits: evolutionary insights from the potato late blight pathogen *Phytophthora infestans*. BMC Evol. Biol. 10:283.
- Montarry, J., Cartier, E., Jacquemond, M., Palloix, A., Moury, B. 2012. Virus adaptation to quantitative plant resistance: erosion or breakdown? J. Evol. Biol. 25:2242–2252.
- Morris, C.E., Moury, B. 2019. Revisiting the concept of host range of plant pathogens. Annu. Rev. Phytopathol. 57:63–90.
- Morris, C.E., Lamichhane, J.R., Nikolić, I., Stanković S., Moury, B. 2019. The overlapping continuum of host range among strains in the *Pseudomonas syringae* complex. Phytopathology Research 1:4.
- Morris, C.E., Glaux, C., Latour, X., Gardan, L., Samson, R., Pitrat, M. 2000. The relationship of host range, physiology and genotype to virulence on cantaloupe in *Pseudomonas syringae* from cantaloupe blight epidemics in France. Phytopathology 90:636-646.
- Morris CE, Sands DC, Vanneste JL, Montarry J, Oakley B, Guilbaud C, Glaux C (2010). Inferring the evolutionary history of the plant pathogen *Pseudomonas syringae* from its biogeography in headwaters of rivers in North America, Europe and New Zealand. mBio 1: e00107-10.
- Newman, M.E.J. 2006. Finding community structure using the eigenvectors of matrices. Phys. Rev. E 74:036104.
- Newman M.E.J., Girvan, M. 2004. Finding and evaluating community structure in networks. Phys. Rev. E 69:026113.
- Nicot, P.C., Bardin, M., Dik, A.J. 2002. Basic methods for epidemiological studies of powdery mildew: Culture and preservation of isolates, production and delivery of inoculum, and disease assessment. Pages 83-99 in: The Powdery Mildews: A Comprehensive Treatise. R. R. Bélanger, W. R. Bushnell, A. J. Dik, and T. L. W. Carver, eds. The American Phytopathological Society, St. Paul, MN.
- Parisi, L., Morgaint, B., Blanco Garcia, J., Guilbaud, C., Chandeysson, C., Bourgeay, J.-F., Moronvalle, A., Brun, L., Brachet M.-L., Morris, C.E. 2019. Bacteria from four phylogroups of the *Pseudomonas syringae* complex can cause bacterial canker of apricot. Plant Pathol. 68:1249-1258.
- Parlevliet, J. E. 1977. Evidence of differential interaction in the polygenic *Hordeum vulgare Puccinia hordei* relation during epidemic development. Phytopathology 67:776-778.
- Patefield, W.M. 1981. Algorithm AS159. An efficient method of generating r x c tables with given row and column totals. J. Royal Stat. Soc. 30:91–97.
- Perchepied, L., Bardin, M., Dogimont, C., Pitrat, M. 2005. Relationship between loci conferring downy mildew and powdery mildew resistance in melon assessed by quantitative trait loci mapping. Phytopathology 95:556-565.
- Peters, A.R., Zhang, Z., Richards, J.K., Friesen, T.L., Faris, J.D. 2019. Genetics of variable disease expression conferred by inverse gene-for-gene interactions in the wheat-*Parastagonospora nodorum* pathosystem. Plant Physiol. 180:420-434.

- Pons, P., Latapy, M. 2006. Computing communities in large networks using random walks. J. Graph Algorithms Appl. 10:191–218.
- Poulicard, N., Pinel-Galzi, A., Hébrard, E., Fargette, D. 2010. Why *Rice yellow mottle virus*, a rapidly evolving RNA plant virus, is not efficient at breaking *rymv1-2* resistance? Mol. Plant Pathol. 11:145-154.
- Quenouille, J., Paulhiac, E., Moury, B., Palloix, A. 2014. Quantitative trait loci from the host genetic background modulate the durability of a resistance gene: a rational basis for sustainable resistance breeding in plants. Heredity 112:579-587.
- Quillévéré-Hamard, A., Le Roy, G., Moussart, A., Baranger, A., Andrivon, D., Pilet-Nayel, M.-L., Le May, C. 2018. Genetic and pathogenicity diversity of *Aphanomyces euteiches* populations from pea-growing regions in France. Front. Plant Sci. 9:1673.
- Quillévéré-Hamard, A., Le Roy, G., Lesné, A., Le May, C., Pilet-Nayel, M.-L. 2020. Aggressiveness of diverse French *Aphanomyces euteiches* isolates on pea Near-Isogenic-Lines differing in resistance QTL. Phytopathology, doi: 10.1094/PHYTO-04-20-0147-R.
- Raghavan, U.N., Albert, R., Kumara, S. 2007. Near linear time algorithm to detect community structures in large-scale networks. Phys. Rev. E 76:036106.
- Reichardt, J., Bornholdt, S. 2006. Statistical mechanics of community detection. Phys. Rev. E 74:016110.
- Sacristán, S., García-Arenal, F. 2008. The evolution of virulence and pathogenicity in plant pathogen populations. Mol. Plant Pathol. 9:369-384.
- Saintenac, C., Lee, W.-S., Cambon, F., Rudd, J.J., King, R.C., Marande, W., Powers, S.J., Bergès, H., Phillips, A.L., Uauy, C., Hammond-Kosack, K.E., Langin, T., Kanyuka, K. 2018. Wheat receptor-kinase-like protein *Stb6* controls gene-for-gene resistance to fungal pathogen *Zymoseptoria tritici*. Nat. Genet. 50:368-374.
- Soltis, N.E., Atwell, S., Shi, G., Fordyce, R., Gwinner, R., Gao, D., Shafi, A., Kliebenstein D.J. 2019. Interactions of tomato and *Botrytis cinerea* genetic diversity: Parsing the contributions of host differentiation, domestication, and pathogen variation. Plant Cell 31:502–519.
- Staniczenko, P.P.A., Kopp, J.C., Allesina, S. 2013. The ghost of nestedness in ecological networks. Nat. Commun. 4:1391.
- Suweis, S., Simini, F., Banavar, J.R., Maritan, A. 2013. Emergence of structural and dynamical properties of ecological mutualistic networks. Nature 500:449–452.
- Tellier, A., Brown, J.K.M. 2011. Spatial heterogeneity, frequency-dependent selection and polymorphism in host-parasite interactions. BMC Evol. Biol. 11:319.
- Thrall, P.H., Barrett, L.G., Dodds, P.N., Burdon, J.J. 2016. Epidemiological and evolutionary outcomes in gene-for-gene and matching allele models. Front. Plant Sci. 6:1084.
- Traag, V.A., Bruggeman, J. 2009. Community detection in networks with positive and negative links. Phys. Rev. E 80:036115.
- Valverde, S., Elena, S.F., Solé, R. 2017. Spatially induced nestedness in a neutral model of phage-bacteria networks. Virus Evol. 3:vex021.
- Valverde, S., Jordi Piñero, J., Corominas-Murtra, B., Montoya, J., Joppa, L., Solé, R. 2018. The architecture of mutualistic networks as an evolutionary spandrel. Nat. Ecol. Evol. 2:94–99.

- Vera Cruz, C.M., Bai, J.F., Ona, I., Leung, H., Nelson, R.J., Mew, T.W., Leach J.E. 2000. Predicting durability of a disease resistance gene based on an assessment of the fitness loss and epidemiological consequences of avirulence gene mutation. Proc. Natl. Acad. Sci. USA 97:13500-13505.
- Wang, M., Roux, F., Bartoli, C., Huard-Chauveau, C., Meyer, C., Lee, H., Roby, D., McPeek, M.S., Bergelson, J. 2018. Two-way mixed-effects methods for joint association analysis using both host and pathogen genomes. Proc. Natl. Acad. Sci. USA 115:E5440-E5449.
- Weitz, J.S., Poisot, T., Meyer, J.R., Flores, C.O., Valverde, S., Sullivan, M.B., Hochberg, M.E. 2013. Phage–bacteria infection networks. Trends Microbiol. 21:82–91.
- Wichmann, G., Bergelson, J. 2004. Effector genes of *Xanthamonas axonopodis* pv. *vesicatoria* promote transmission and enhance other fitness traits in the field. Genetics 166:693-706.
- Zhong, Z., Marcel, T.C., Hartmann, F.E., Ma, X., Plissonneau, C., Zala, M., Ducasse, A., Confais, J., Compain, J., Lapalu, N., Amselem, J., McDonald, B.A., Croll, D., Palma-Guerrero, J.C. 2017. A small secreted protein in *Zymoseptoria tritici* is responsible for avirulence on wheat cultivars carrying the *Stb6* resistance gene. New Phytol. 214:619-631.