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#### Integrated control of potato late blight: predicting the combined efficacy of host resistance and fungicides

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## Integrated control of potato late blight: predicting the combined efficacy of host resistance and fungicides

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1	Short Title: Predicting combined efficacy
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3 4	Integrated control of potato late blight: predicting the combined efficacy of host resistance and fungicides
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17	

#### 18 Abstract

19 Integrating cultivars that are partially resistant with reduced fungicide doses offers growers an opportunity to decrease fungicide input but still maintain disease control. To use integrated 20 21 control strategies in practice requires a method to determine the combined effectiveness of 22 particular cultivar and fungicide dose combinations. Simple models, such as additive dose 23 models (ADM) and multiplicative survival models (MSM), have been used previously to 24 determine the joint action of two or more pesticides. This study tests whether a model based 25 on multiplicative survival principles can predict the joint action of fungicide doses combined with varieties of differing partial host resistance. Data from eight field experiments on potato 26 27 late blight (*Phytophthora infestans*), where the severity of foliar blight was assessed and 28 converted to AUDPC, were used to test the model. A subset of data, derived from the most 29 susceptible cultivar, King Edward, was used to produce dose response curves from which 30 parameter values were estimated, quantifying fungicide efficacy. These values, along with the 31 untreated values for the more resistant cultivars, Cara and Sarpo Mira, were used to predict 32 the combined efficacy of the remaining cultivar by fungicide dose combinations. Predicted efficacy was compared against observations from an independent sub-set of treatments from 33 34 the field experiments. The analysis demonstrated that multiplicative survival principles can 35 be applied to describe the joint efficacy of host resistance and fungicide dose combinations.

#### 36 Introduction

There are many ways in which integrated control strategies for foliar plant pathogens can be 37 deployed. Examples of using host resistance to limit damage or reduce dependence on 38 39 fungicides include switching from susceptible to moderately resistant varieties for control of 40 northern corn leaf blight of maize (Exserohilum turicum) and decreasing the number of fungicide applications on wheat varieties with resistance to tan spot (Drechslera tritici-41 42 repentis) (Debela et al., 2017, Jørgensen & Olsen, 2007). Decreasing fungicide inputs on moderately resistant potato cultivars compared with susceptible cultivars has been shown to 43 be an effective control strategy against late blight, caused by *Phytophthora infestans* (Fry, 44 45 1978, Gans et al., 1995, Nærstad et al., 2007). Despite considerable research demonstrating 46 the potential to optimise inputs by combining strategies, e.g. fungicide inputs and genetic 47 resistance of varieties, no study has yet determined whether there is a predictable relationship describing the joint action of different components of integrated control. 48

49 Simple models to predict the joint action of two or more pesticides applied in mixture have been used in laboratory and field studies for invertebrate pests, weeds and diseases 50 51 (Bliss, 1939, Scardavi, 1966, Colby, 1967, Rummens, 1975, Gisi et al., 1985, Paveley et al., 52 2003). In these simple models, the efficacy of each component of the mixture is quantified 53 and used to predict the efficacy of the mixture. The efficacy of treatment combinations has 54 been described as synergistic or antagonistic where the joint action of the mixture 55 components exceeded or failed to achieve the level of control predicted. More recently, one such model has been applied to assess the joint action of host resistance genes (identified as 56 57 quantitative trait loci) against diseases of winter wheat (Grimmer et al., 2015).

58 The two most frequently used models to determine the combined efficacy of two or 59 more control methods have been defined as two broad types, additive dose models (ADM)

and the multiplicative survival models (MSM), also known as the Abbot and Wadley 60 61 methods respectively. Their respective appropriateness for joint action comparisons has been reviewed previously (Morse, 1978). The ADM assumes that the action of one component can 62 63 be directly substituted for the action of the other, i.e. that the dose of one mixture component can be expressed as an equivalent dose of the other component. It has been reported 64 65 previously that the effects of host resistance and fungicides to control *Phytophthora infestans* 66 were 'additive' (Fry, 1978). However, the assumption underlying the ADM cannot be met 67 when the two components have fundamentally different modes of action, such as in the case described here where host resistance and fungicides are combined. MSM, however, calculates 68 69 the proportion of the pathogen population which 'survives' the effect of each component 70 separately, and then predicts the proportion of the population which would survive joint use 71 of the control methods by multiplying the survivorship proportions. The underlying 72 assumption is that the two components act independently, which is a plausible assumption for 73 host resistance and fungicides. Hence, MSM is likely to be the more appropriate model for 74 determining the joint action of host resistance and fungicide dose.

This study tested whether a simple multiplicative survival model can predict the joint action of fungicide dose and host resistance against *Phytophthora infestans* on potato and could provide an accurate prediction of the performance of those combinations under field conditions.

79

#### 80 Materials and methods

Data derived from eight integrated control field experiments conducted in 2010 and 2011 were used in this study. The experiments were conducted on late blight (*Phytophthora infestans*) of potato (*Solanum tuberosum*) at two sites: Ayrshire and Ceredigion in the UK.

#### 4

Two experiments were conducted at each site in each year: one with treatments applied during rapid canopy growth ('rapid canopy') and the other with treatments applied later in the seasons when canopy size was relatively stable ('stable canopy').

87

#### 88 Experimental design

89 Treatments in each experiment consisted of all combinations of three cultivars and four 90 fungicide doses, with an untreated control for each cultivar (Table 1). Integrated control 91 treatments were considered to be the treatments where cultivars, which were moderately 92 (Cara) or highly (Sarpo Mira) resistant, were combined with fungicide doses below the 93 maximum permitted dose per application (the full label recommended dose). At both sites 94 experiments were laid out as a randomised split-plot design with four replicates for each 95 treatment. Fungicides were applied at the whole plot level and cultivars planted at the subplot level. Fungicides were randomised within each block and varieties randomised within each 96 97 fungicide plot. In Ceredigion, main plots were four rows wide (each row = 0.9m wide) by 98 11m long. Main plots were separated by 1.5 m unplanted row length. Each main plot was divided into cultivar sub-plots of four rows by 3.0m long and separated by 1.0m unplanted 99 100 row length. All cultivars were supplied as the same seed size (35 to 45mm) and planted at 101 30cm spacing. A single row of King Edward was planted between each of the blocks as an 102 infector row.

In Ayrshire in 2010, fungicide treatment plots were four rows wide by 9.75 m long and separated longitudinally by 1.5 m of bare earth. Each cultivar sub-plot was four rows by 2.75 m long separated by 0.75 m unplanted row length with 25cm seed spacing. In 2011, seed spacing was 0.23 m. The fungicide treatment plots were 8.97 m long and the cultivar plots 2.3

m long. The unplanted row length was 1.61 m. A single row of King Edward was planted
longitudinally between each of the blocks as an infector row.

One fungicide, mandipropamid [full recommended label rate 0.6 l ha<sup>-1</sup> (250g l<sup>-1</sup>) as 109 Revus, Syngenta Ltd] was applied to cultivars at a range of doses, as proportions of the full 110 111 recommended label rate (Table 1). Treatments were applied either during rapid canopy and or during stable canopy growth in separate experiments. For the rapid canopy experiments, first 112 113 treatment fungicides were applied at the time of the first blight warning or when plants met within the rows, whichever was soonest. For the stable canopy experiments, chlorothalonil + 114 propamocarb-hydrochloride [full recommended rate 2.5 l ha<sup>-1</sup> (375 g l<sup>-1</sup> + 375 g l<sup>-1</sup>) as Merlin, 115 Bayer CropScience] were applied to all plots at 7- or 10-day intervals until rapid canopy 116 117 growth was complete (typically three applications at 10-day intervals per season). For all 118 experiments, four fungicide applications for each treatment at 7-day intervals were planned, however, there was flexibility depending on the epidemic progress at different sites and for 119 120 different seasons (Table 5). In Ceredigion, there were four applications of treatment 121 fungicides in app experiments conducted in 2010 and 2011. In Ayrshire, there were four test 122 fungicide applications in the 2010 rapid canopy experiment and five in the equivalent trial in 2011. Both stable canopy trials had six applications of test fungicides. Once treatment sprays 123 had been applied, all plots were sprayed with between 1274g ha<sup>-1</sup> and 1540g ha<sup>-1</sup> mancozeb 124 (as Dithane NT, Penncozeb or Laminator Flo depending on site and season) at 7-day intervals 125 126 until desiccation. In Ceredigion, fungicide treatments were applied using a handheld Oxford 127 Precision Sprayer in 250 litres of water per hectare operating at 200 kPa through 110° flat fan 128 nozzles. In Ayrshire, treatments were applied in 200 litres of water per hectare, using a 129 tractor-mounted modified AZO compressed air sprayer through Lurmark F03-110 flat fan 130 nozzles at an operating pressure of 350 kPa.

131	Infector rows (cv. King Edward) were not sprayed with fungicide and were inoculated
132	with isolate(s) of genotype 13_A2 grown on Rye B agar (Caten & Jinks, 1968) but sub-
133	cultured several times on detached King Edward potato leaves prior to inoculation of
134	experiments. These isolates were used to produce a sporangial suspension (a minimum of 1 x
135	10 <sup>4</sup> spores ml <sup>-1</sup> ) in sterile distilled water which was applied to the spreader rows using a
136	handheld mister. The isolates were supplied by the James Hutton Institute, Dundee, UK
137	(Table 2).

Foliar blight was assessed at least weekly as the percentage of leaf area affected by *P*. *infestans*, with more frequent assessments when the epidemic was increasing rapidly, using the modified MAFF key 2.1.1: Potato Blight on the Haulm (Anon, 1976; Large, 1952).

141

#### 142 Determining the effectiveness of cultivar and fungicide dose combinations

For each treatment the Area Under the Disease Progress Curve (AUDPC) was calculated from the foliar late blight severity scores (Campbell & Madden, 1990). To predict the effectiveness of host resistance and fungicide combinations, a multiplicative survival equation was derived incorporating a previously published exponential equation (1) describing the fungicide dose response curve, where  $D_d$  is disease severity at dose d and  $D_o$  is disease when fungicide dose = 0 (Paveley *et al.*, 2000).

149 (1) 
$$D_d = D_o [1 - b(1 - e^{-kdose})]$$

150

Data generated from the observed dose response curve of the most susceptible cultivar, King Edward, were used to calculate the parameters b and k, where b represents the amount of disease that might be potentially controlled with an infinite dose and k defines the rate of change of disease severity with dose. Curves were forced through the untreated values and

parameters calculated using the following equation in FITNONLINEAR in Genstat 16<sup>th</sup> 155 156 edition (VSN International Ltd, UK).

157

The dose response equation and parameters were then used in the following equation (2) to 158 predict the effectiveness of cultivar and fungicide dose combinations based on the principles 159 160 of multiplicative survival:

(2) 
$$D = D_o \left[ \left( \frac{D_r}{D_s} \right) (1 - b(1 - e^{-kdose})) \right]$$

161 D is the predicted level of disease for the appropriate cultivar and fungicide dose combination,  $D_0$  is the untreated AUDPC of the standard susceptible cultivar (in this case the 162 most susceptible cultivar King Edward). For the first analysis,  $D_s$  is the untreated AUDPC for 163 164 the standard cultivar,  $D_r$  is the untreated AUDPC for the partially resistant test cultivar and 165 *dose* is the proportion of the full fungicide dose.

AUDPC values were logit transformed using an equation (3) modified from Grimmer 166 (M-s)] 167 *et al.* (2015):

168 (3) 
$$LTS = In[s/(M-s)]$$

Where LTS is the logit transformed AUDPC, In is the natural logarithm, s is the observed or 169 170 predicted disease severity and M is the maximum AUDPC achievable during the disease assessment period (e.g. if disease assessment period was 59 days then maximum AUDPC is 171 172 5900). The transformed observed severity was linearly regressed against the transformed predicted severity as advocated in Piñeiro et al. (2008). All analysis was done in Genstat 16<sup>th</sup> 173 174 Edition (VSN International Ltd, UK).

Results 175

176 Parameter estimates for the dose response curves were derived from dose response curves generated using King Edward as the baseline. The fitted dose response curves were close to 177 the observed AUDPC values (Figure 1). The percentage variance accounted for (as  $R^2$ ) for all 178 fitted curves ranged from 96% to 100% (Table 3). Observed and predicted disease severities, 179 as the AUDPC for each cultivar and fungicide dose combination, for moderately resistant 180 181 Cara and highly resistant Sarpo Mira cultivars were compared (Table 4). There was a highly significant relationship (P < 0.001;  $R^2 = 0.88$ ) between the predicted and observed values for 182 each fungicide dose and cultivar combination (Figure 2). When the combined effects of 183 184 fungicide dose and cultivar were predicted, the resultant AUDPC values were generally 185 higher than those observed in the experiments, regardless of cultivar resistance rating.

186 The percentage of foliar late blight present at the first and last fungicide application in 187 each trial varied depending on the site and year (Table 5). In 2010, only the stable canopy 188 trial in Ayrshire had foliar late blight present when first fungicides were applied. In contrast 189 in 2011, most King Edward and Cara treatments had traces of foliar late blight when first fungicides were applied. At the time the final fungicide application was applied to King 190 191 Edward, foliar late blight ranged from 0.7 to 94% leaf area affected and 5 out of the 8 trials 192 had >90% foliar late blight. In comparison, for Cara, foliar late blight ranged from 0.5 to 85% and for Sarpo Mira 0.03 to 7.8% leaf area affected. Conditions were generally not 193 194 favourable for disease development early in the season at the Ayrshire site in 2010. The 195 epidemic was slower to start at the Ceredigion site in the rapid canopy trials in both 2010 and 196 2011.

Following the logit transformation, which took into account the differences between experiments in the duration of the disease assessment period (and hence differences in the maximum possible AUDPC; equation 3), the regression accounted for 76% of the variation in the logit transformed severity [P=<0.001, slope and intercept 95% confidence intervals

- 201 (0.7144, -1.569) and (0.5341, -2.324)] (Figure 3). T-values [36.01, -15.59] demonstrated that
- this line was significantly different from the slope of 1 and intercept of 0.

203

to per period

#### 204 **Discussion**

The analysis presented here shows that multiplicative survival principles can be 205 applied to derive a simple model to describe the efficacy of host resistance and fungicide 206 207 dose combinations. To generate predicted AUDPC values and test the ability of the model to 208 predict the joint action of host resistance and fungicide dose, there were two requirements: a 209 fungicide dose response curve (including an 'untreated' control) for a susceptible cultivar (in 210 this case King Edward) plus an 'untreated' control for each test cultivar. Using this method, 211 there was a good relationship between observed AUDPC values and the multiplicative 212 survival model (MSM) predictions.

213 In a previous study, dose response curves for different host resistance/fungicide dose 214 treatments were compared and differences used to identify the contribution of host resistance. 215 The contribution of host resistance was expressed as the equivalent dose of fungicide 216 required to match the additional disease control provided by a more resistant cultivar (Fry, 217 1978). An alternative approach, using three dimensional regression, was used to estimate the 218 equivalent fungicide dose equivalent to one point on a 1 to 9 scale (where 9 is most resistant) 219 which defined cultivar resistance (Gans et al., 1995). Such approaches have the disadvantage 220 of requiring dose response curves from all treatments to compare and quantify the benefits of 221 host resistance. This paper demonstrates, for the first time, a method in which the 222 performance of a cultivar in combination with different fungicide doses, could be predicted, 223 without the need to include all the fungicide dose combinations in the experiment and in the 224 absence of a cultivar resistance rating.

MSM models were used originally to determine whether the joint action of mixture components was synergistic. Experimental results are compared with the reference model which represents the joint action predicted from the efficacy of the components. Where the observed severity or AUDPC values were less than, or more than, the predicted values, the

combination may be considered synergistic or antagonistic, respectively (Kosman & Cohen, 1996). For the data presented in this paper, the majority of observed values were below their corresponding predicted values. Cultivar and fungicide combinations therefore performed better than predicted for the majority of host resistance and fungicide dose combinations. Although this is a positive outcome for the value of integrated control, it is important to consider whether this apparent synergy between host resistance and fungicide treatment may in fact have been an artifact of the experimental method.

236 As described in the methods, mancozeb was applied to all treatments, including 237 'untreated' plots, once treatment fungicide applications were completed. This over-spray was 238 designed to allow more time for differences in foliar late blight between treatments to 239 develop prior to defoliation. By the time the final treatment fungicides were applied, >90% of 240 the leaf area in untreated King Edward plots, in five out of eight of the experiments, was 241 infected with P. infestans. Mancozeb prevents spore gemination, but has limited effects on 242 established infections and mycelial growth (Bruck et al., 1981, Kaars Sijpesteijn, 1982). 243 Mancozeb application where > 0.5% of leaf area was affected by *P. infestans* has been shown 244 previously to be insufficient to decrease epidemic growth rate immediately, with a delay of 8 245 to 10 days before established epidemics were slowed (Fry et al., 1979). It is likely, therefore, 246 that the timing of mancozeb application, relative to epidemic severity and growth rate, 247 differed depending on cultivar and the fungicide dose applied in the experimental treatments. 248 It has been demonstrated previously that the order in which fungicides are applied can also 249 impact on the ability of particular fungicide products to influence the epidemic (Bain & 250 Bardsley, 2009). In the current study, the effect of mancozeb on the epidemic growth rate was 251 likely to be lower for treatments where disease was well established (e.g. on untreated King 252 Edward) compared with other treatments where disease was less established at the time of 253 mancozeb application (e.g. on untreated Sarpo Mira) leading to bias. Such bias could not be

excluded from this analysis, given the absence of a completely fungicide untreated control. Given the apparent effect of oversprays on the epidemic and the potential for bias, it is suggested that such over-sprays should be avoided, or completely untreated controls should be included, for future experiments.

Generating up to date information on the likely efficacy of host resistance and 258 259 fungicide dose combinations is necessary, particularly when pathogen populations are 260 evolving rapidly towards aggressiveness, virulence or fungicide insensitivity. In Great 261 Britain, the dominance of 13 A2, one of the newer aggressive and more virulent genotypes, 262 resulted in the re-grading of cultivars, including Cara which was used in this experiment, 263 from highly resistant to moderately resistant (Lees et al., 2012). Similarly P. infestans 264 genotypes that are less sensitive to and less well controlled by the fungicide fluazinam have 265 been detected in Europe recently (Schepers, 2017). It has been demonstrated previously that 266 the rank order of partial resistance of cultivars exposed to *P. infestans* remains similar with 267 and without fungicide treatment (Bain et al., 2014), therefore using multiplicative survival 268 principles to explore the potential for using integrated control in the way described here 269 should provide a useful guide to the performance of integrated host resistance and fungicide 270 strategies.

Achieving effective control of late blight on potato using decreased fungicide doses 271 272 on moderately resistant cultivars has been demonstrated previously (Fry, 1975, Clayton & 273 Shattock, 1995, Gans et al., 1995, Bain et al., 2014) and, in some instances, such information 274 has been incorporated into models to guide fungicide applications (Nærstad et al., 2007, Liu 275 et al., 2017). For potato late blight, foliar resistance ratings are calculated for cultivars in 276 many countries (e.g. AHDB, 2017). For the cultivar resistance ratings reported in Europe, 1 277 to 9 ratings are based on AUDPC values for untreated test cultivars, expressed relative to 278 AUDPC values for one susceptible and one resistant reference cultivar. Cultivars with the

same 1 to 9 resistance rating will therefore have very similar relative mean AUDPC values.
Consequently, one resistance rating in the equation should also give a useful indication of the
combined efficacy of fungicide doses on all cultivars with the same rating, e.g. there are
currently 11 cultivars in Great Britain rated 7 for foliar blight resistance.

The method defined in Equation 2 should be more generally applicable both for 283 284 potato late blight worldwide and for integrated control in other pathosystems – although 285 further experimental proof is required. The experimental data required to estimate parameters 286 is relatively simple to obtain and analysis is straightforward. Models to describe the joint 287 action of fungicides mixtures, based on MSM principles, have been included as a component 288 in decision support system models for winter wheat disease control (Paveley et al., 2003, 289 Milne et al., 2007) and models of pathogen evolution which consider integrated control 290 (Carolan et al., 2017). MSM principles have proved to be remarkably generalisable in their 291 application, provided the control methods for which joint action is being calculated are 292 reasonably independent. The principles have proved useful for predicting joint action of 293 fungicides, herbicides and insecticides, joint action of separate fungicide treatments in a spray 294 programme (Paveley et al., 2003), joint action of host resistance QTL (Grimmer et al., 2015) 295 and now for the joint action of host resistance and fungicides.

296

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301	References
302	
303	AHDB, 2017. Potato variety database (Great Britain) [http://varieties.ahdb.org.uk/] accessed
304	7 June 2017.
305	Anon, 1976. Manual plant growth stages and disease assessment keys. MAFF Publications,
306	Middlesex, UK.
307	Bain R, Bardsley E, 2009. The impact of fungicide spray order on foliar blight control in two
308	growing seasons. In: Schepers HTAM, ed. Proceedings of the Eleventh Euroblight
309	Workshop, Hamar, Norway. PPO Special Report 13. Wageningen, The Netherlands, 115-
310	122.

Bain R, Ritchie F, Lees A, Dyer C, 2014. Impact of fungicide input on leaf blight 311 (Phytophthora infestans) development on different cultivars. In: Schepers, HTAM, ed. 312 Proceedings of the Fourteenth Euroblight Workshop. PPO Special Report No. 16. 313 Wageningen, The Netherlands, 65-73. 314

- Bliss C, 1939. The toxicity of poisons applied jointly. Annals of Applied Biology 26, 585-615. 315
- Bruck R, Fry W, Apple A, Mundt C, 1981. Effect of protectant fungicides on the 316 317 developmental stages of *Phytophthora infestans* in potato foliage. *Phytopathology* 71, 164-6.
- 318 Campbell C, Madden L, 1990. Introduction to plant disease epidemiology. New York, USA: John Wiley & Sons, Inc. 319
- 320 Carolan K, Helps J, Van Den Berg F, Bain R, Paveley N, Van Den Bosch F, 2017. Extending the durability of cultivar resistance by limiting epidemic growth rates. Proceedings of the 321 Royal Society B 284, 20170828. 322
- 323 Caten C, Jinks J, 1968. Spontaneous variability of single isolates of *Phytophthora infestans*. I.
- 324 Culture variation. Canadian Jouranl of Botany 46, 329-48.

- 325 Clayton R, Shattock R, 1995. Reduced fungicide inputs to control *Phytophthora infestans* in
- potato cultivars with high levels of polygenic resistance. *Potato Research* **38**, 399-405.
- Colby S, 1967. Calculating synergistic and antagonistic responses of herbicide combinations. *Weed Science* 15, 20-2.
- Debela M, Dejene M, Abera W, 2017. Management of Turicum leaf blight [*Exserohilum turicum* (Pass.) Leonard & Suggs] of maize (*Zea Mays* L.) through integration of host resistance and fungicide at Bako, Western Ethiopia. *African Journal of Plant Science* **11**, 6-322.
- Fry W, 1975. Integrated effects of polygenic resistance and a protective fungicide on development of potato late blight. *Phytopathology* **65**, 908-11.
- Fry W, 1978. Quantification of general resistance of potato cultivars and fungicide effects for
  integrated control of potato late blight. *Phytopathology* 68, 1650-5.
- Fry W, Bruck R, Mundt C, 1979. Retardation of potato late blight epidemics by fungicides
  with eradicant and protectant properties. *Plant Disease Reporter* 63, 970-4.
- 339 Gans P, Carson W, N P, Owen L, 1995. Exploiting cultivar resistance to control potato blight
- 340 (*Phytophthora infestans*). In: Dowley LJ, Bannon E, Cooke L, Keane T, O'sullivan E, eds.
- 341 *Phytophthora infestans 150.* Dublin, Ireland: Boole Press Ltd, 345-50.
- Gisi U, Binder H, Rimbach E, 1985. Synergistic interactions of fungicides with different modes of action. *Transactions of the British Mycological Society* **85**, 299-306.
- 344 Grimmer M, Boyd L, Clarke S, Nd P, 2015. Pyrimiding of partial disease resistance genes
- has a predictable, but diminishing, benefit to efficacy. *Plant Pathology* **64**, 748-53.
- Jørgensen L, Olsen L, 2007. Control of tan spot (*Drechslera tritici-repentis*) using cultivar
- resistance, tillage methods and fungicides. *Crop Protection* **26**, 1606-16.

- 348 Kaars Sijpesteijn A, 1982. Mechanism of action of fungicides. In. Dekker J, Georgopoulos S,
- 349 eds. Fungicide Resistance in Crop Protection. Centre for Agricultural Publishing and
- 350 Documentation, Wageningen, The Netherlands, 32-45.
- Kosman E, Cohen Y, 1996. Procedures for calculating and differentiating synergism and antagonism in action of fungicide mixtures. *Phytopathology* **86**, 1263-72.
- Large EC, 1952. The interpretation of progress curves for potato blight and other plant diseases. *Plant Pathology* **1**, 109-117.
- Lees A, Stewart J, Lynott J, Carnegie S, Campbell H, Roberts A, 2012. The effect of a dominant *Phytophthora infestans* genotype (13\_A2) in Great Britain on host resistance to foliar late blight in commercial potato cultivars. *Potato Research* **55**, 125-34.
- Liu Y, Langemeier M, Small I, Joseph L, Fry W, 2017. Risk management strategies using precision agriculture technology to manage potato late blight. *Agronomy Journal* **109**, 562-75.
- Milne A, Paveley N, Audsley E, Parsons D, 2007. A model of the effect of fungicides on
  disease-induced yield loss, for use in wheat disease management decision support systems. *Annals of Applied Biology* 151, 113-25.
- Morse PM, 1978. Some comments on the assessment of joint action of herbicide mixtures. *Weed Science* 26, 58-71.
- Nærstad R, Hermansen A, Bjor T, 2007. Exploiting host resistance to reduce use of fungicides to control potato late blight. *Plant Pathology* **56**, 156-66.
- Paveley N, Lockley D, Vaughan T, Thomas J, Schmidt K, 2000. Predicting effective doses
  through observation of leaf emergence. *Plant Pathology* 49, 748-66.
- Paveley N, Thomas J, Vaughan T, Havis N, Jones D, 2003. Predicting effective doses for the
  joint action of two fungicide applications. *Plant Pathology* 52, 638-47.

Piñeiro G, Perelman S, Guerschman J, Paruelo J, 2008. How to evaluate models: observed vs.

predicted or predicted vs. observed? *Ecological Modelling* **216**, 316-22.

Rummens F, 1975. An improved definition of synergistic and antagonistic effects. *Weed Science* 23, 4-6.

376 Scardavi A, 1966. Synergism among fungicides. *Annual Review of Phytopathology* 4, 335-48.

377

Schepers HTAM, 2017. Reduced sensitivity to fluazinam detected in the rapidly increasing *Phytophthora infestans* lineage EU-37 (Dark green 33). [goo.gl/cmCw3p]. Accessed 12

380 February 2018.

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#### 382 Figures

Figure 1. Dose response curves (lines) derived from data in the rapid canopy experiments and fitted to the original AUDPC values (points) for King Edward using the equation (1) derived from Paveley *et al.*, 2000. Grey lines are the Ayrshire site and black lines the Ceredigion site. Solid lines are data from 2010 and dashed lines are data from 2011.

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Figure 2. Relationship between observed and predicted disease severity for Cara (circles) and
Sarpo Mira (triangles) from rapid and stable canopy trials conducted in 2010 and 2011. Black
data points identify data points from Ceredigion and white data points from Ayrshire.

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Figure 3. Relationship between logit transformed observed and predicted disease severity for Cara (circles) and Sarpo Mira (triangles) from rapid and stable canopy trials conducted in 2010 and 2011. Slope and intercept 95% confidence intervals [(0.7144, -1.569) and (0.5341, -

395	2.324)]. Solid black line represents the regression and dotted line the 1:1 line. Black data
396	points identify data points from the Ceredigion site and white data points from the Ayrshire
397	site.

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### 1 Figures

- 2 Figure 1.
- 3



6 Figure 2







#### Tables

Table 1. Cultivars, fungicide treatments, spray interval and UK foliar blight resistance ratings

for all varieties included in the rapid canopy and stable canopy experiments.

Year	Fungicide doses applied (as percentage of full recommended label rate)	Spray interval	Cultivar (foliar blight resistance rating)*			
2010 and 2011	0, 25%, 50%, 75% and 100%	7 days	King Edward (3)	Cara (5)	Sarpo Mira (7)	

\*from the Potato Variety Database (AHDB, 2017).

Table 2. P. infestans isolate(s) and inoculation dates for each site by year.

			Site			
	Ayrsh	ire	Ceredigion			
Year	inoculation date(s)**	isolates	inoculation date	isolates		
2010	15, 19, 28 July 17 August	2009_7654A 07/39, 2009_7654A,	3 July	2009_7654A		
2011	8, 13, 18, 28 July 8, 16 August	2006_3928A, 2008_6082F	12 July	2009_7654A		

\*\* There were multiple trials at the site, inoculated on different dates.

Table 3. Untreated AUDPC values for King Edward, Cara and Sarpo Mira plus parameter estimates (refer to text for definitions) derived from dose response curves (cv. King Edward/fungicide dose) and  $R^2$  for the fitted dose response curves by site, experiment and year.

			Untreated AUDPC			Parameter		
Experiment	Site <sup>a</sup>	Year	King Edward	Cara	Sarpo Mira	b	k	$R^2$
Rapid canopy	AYR	2010	1643	196	22	0.94	19.40	1.00
		2011	4008	1957	431	0.81	8.17	1.00
	CER	2010	3741	3042	786	0.81	10.73	1.00
		2011	2686	2058	71	0.86	7.10	0.99
Stable canopy	AYR	2010	3112	1371	36	0.76	6.10	1.00
		2011	2419	1568	116	0.75	9.42	1.00
	CER	2010	3828	3234	918	0.40	3.97	0.96
		2011	2094	1885	204	0.56	4.28	0.98

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<sup>a</sup>AYR = Ayrshire site, CER = Ceredigion site.

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Table 4. Observed and predicted disease severity (as AUDPC) of foliar blight for rapid canopy and stable canopy experiments. Fungicides were applied to rapid canopy experiments early in the season when plants are actively growing and once canopy expansion was complete for stable canopy experiments. Dose is expressed as a proportion of the maximum permitted dose per application and cultivar is expressed as a 1 to 9 ranking, where 1 is most susceptible and 9 is most resistant to *P. infestans*.

		Cultivar			Т	rial	
		(resistance		Rapid C	Canopy	Stable	Canopy
Site	Year	rating)	Dose	Observed	Predicted	Observed	Predicted
Ayrshire	2010	Cara (5)	0.25	23	12	234	557
2			0.5	24	11	45	380
			0.75	22	11	48	341
			1.0	21	11	35	333
		Sarpo Mira (7)	0.25	13	1	21	15
			0.5	13	1	17	10
			0.75	10	1	16	9
			1.0	15	1	15	9
Ayrshire	2011	Cara (5)	0.25	431	572	399	510
			0.5	216	392	239	409
			0.75	176	369	264	400
			1.0	136	366	213	399
		Sarpo Mira (7)	0.25	67	133	24	38
			0.5	50	91	19	30
			0.75	49	86	17	30
			1.0	43	85	18	30
Ceredigion	2010	Cara (5)	0.25	167	742	1180	2427
			0.5	115	584	1152	2129
			0.75	133	573	814	2018
			1.0	96	572	658	1977
		Sarpo Mira (7)	0.25	57	192	309	689
			0.5	66	151	305	604
			0.75	66	148	309	573
			1.0	41	148	377	561
Ceredigion	2011	Cara (5)	0.25	203	592	883	1195
			0.5	156	344	607	958
			0.75	105	302	674	877
			1.0	85	294	411	849
		Sarpo Mira (7)	0.25	10	20	44	129
			0.5	6	12	22	103
			0.75	6	10	17	95
			1.0	7	10	22	92

Table 5. The percentage leaf area affected by foliar late blight (%) in untreated varieties at the first (First) or within 7 days of the last (Last) treatment application and the number of test fungicides applied to each trial.

			Perce	ntage of l	eaf area	No. of	No. of			
Experi-					(%	6)			treatment	mancozeb
ment	Site <sup>a</sup>	Year	King E	Edward	Cara Sarpo Mira		applications	applications		
			First	Last	First	Last	First	Last		
Rapid	AYR	2010	0.0	0.7	0.0	0.5	0.0	0.3	4	6
canopy		2011	5.3	100.0	1.3	47.5	0.5	7.3	5	4
	CER	2010	0.0	17.5	0.0	22.0	0.0	0.3	4	8
		2011	0.0	9.0	0.0	0.7	0.0	0.0*	4	4
Stable	AYR	2010	0.2	95.0	0.1	33.0	0.0	1.0	6	2
canopy		2011	0.1	97.0	0.0*	60.0	0.0	4.3	5	2
	CER	2010	0.0	90.0	0.0	62.5	0.0	7.8	4	5
		2011	0.1	93.8	0.1	85.0	0.0	7.5	4	2

<sup>a</sup>AYR = Ayrshire site, CER = Ceredigion site. \* = 0.03

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