

# Population dynamics of the potato cyst nematode, *Globodera pallida* in relation to temperature, potato cultivar and nematicide application

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1 **Population dynamics of the potato cyst nematode, *Globodera***  
2 ***pallida*, in relation to temperature, potato cultivar and nematicide**  
3 **application**

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8

9 The impact of increasing temperatures on the population dynamics of the soil-dwelling  
10 nematode *Globodera pallida*, a persistent and economically important pest of potatoes, was  
11 investigated. The reproductive factor (final population/initial population) and length of life  
12 cycle were found to be temperature sensitive. Pot experiments performed over 4 months  
13 allowed comparison of the effect on development of *G. pallida* of two temperature regimes:  
14 an average temperature comparable to current field conditions (14.3 °C) and an average  
15 temperature above current field conditions (17.3 °C). A larger second generation of juveniles  
16 was observed at 17.3 °C compared to 14.3 °C. Multiplication of *G. pallida* at field sites in  
17 Shropshire and East Lothian (average soil temperatures of 14.1 and 15.1 °C, respectively,  
18 during potato cropping) was also examined. A quantitative PCR assay and visual examination  
19 of roots were used to monitor the dynamics of the *G. pallida* populations in both field sites at  
20 4-weekly intervals. Four cultivars, Desirée, Cara, Maris Piper and Estima, were grown with  
21 and without nematicide treatments. Nematicide treatments suppressed population increases at  
22 both sites. Females were observed on the roots of cvs Cara and Desirée at the end of the

23 growing season in Shropshire, but not at East Lothian, and are likely to represent a second  
24 generation.

25 *Keywords:* population dynamics, potato cyst nematode, second generation, temperature,  
26 nematicide treatment

27

## 28 Introduction

29 The potato cyst nematodes (PCN) *Globodera rostochiensis* and *Globodera pallida* are major  
30 parasites of potatoes and other members of the Solanaceae family. Infected plants typically  
31 have a smaller root system (Trudgill & Cotes, 1983), have a diminished capacity to take up  
32 nutrients from the soil and are adversely affected by water stress or disturbance of nutrient  
33 metabolism. According to the EPPO Global Database (2019), *G. rostochiensis* and *G. pallida*  
34 have been detected in 72 and 49 countries respectively. In the UK, the direct and indirect crop  
35 losses caused by PCN were valued at 9% of yield annually (Evans, 1993) and the economic  
36 cost of PCN to the UK processing and fresh market potato industry was estimated at c. £26  
37 million in 2009 (Twining *et al.*, 2009). Increasing management costs after implementation of  
38 the PCN Directive (2007/33/EU), which requires improved pre-planting soil testing for all  
39 seed crops and an annual survey of 0.5% of ware land, also add to the economic impact of  
40 PCN. In fields where PCN are detected, the directive prohibits the growing of seed potatoes,  
41 and ware potatoes may only be grown under an officially approved control programme that  
42 includes using resistant varieties, nematicides or other control measures (Hockland *et al.*,  
43 2000).

44 The population dynamics of *G. pallida* and *G. rostochiensis* and associated yield loss  
45 of the potato crop differ greatly among years and locations (Greco *et al.*, 1982; Seinhorst,

46 1982). As mentioned above, damage caused by PCN, particularly *G. rostochiensis*, can be  
47 minimized by the selection of resistant potato cultivars within a rotation cycle (Trudgill,  
48 1986). However, with susceptible cultivars, factors including the amount of the initial  
49 population of PCN (Trudgill *et al.*, 2014), environmental factors (Jones, 1983) and soil type  
50 (Trudgill, 1986; Elston *et al.*, 1991) can affect multiplication rates. In general, the larger the  
51 population of *Globodera* spp. in the field at planting, the greater the yield losses (Trudgill *et*  
52 *al.*, 2014; Seinhorst, 1982), although it is also known that PCN reproduction is density  
53 dependent (Trudgill *et al.*, 2014) and at lower densities the multiplication rate is higher.

54         In the last century, global temperatures have been increasing due to climate change  
55 and the effect of this trend on crop losses due to plant pests and diseases, including soil  
56 dwelling organisms such as PCN, is of concern (Ellenby & Smith, 1975; Stanton & Sartori,  
57 1990; Munir *et al.*, 2009, Ebrahimi *et al.*, 2014; Jones *et al.*, 2017; Skelsey *et al.*, 2018).  
58 Recently Jones *et al.* (2017) and Skelsey *et al.* (2018) have reported on the potential impact  
59 of climate change in the UK on the potato cyst nematodes *G. rostochiensis* and *G. pallida*  
60 based on experiments conducted in controlled environments. While the reports differ in some  
61 respects, they generally agree that increases in soil temperatures from current levels are likely  
62 to lead to increased multiplication of PCN, particularly *G. rostochiensis*. *Globodera pallida* is  
63 more adapted to cooler temperatures than *G. rostochiensis* and, hence, increases in soil  
64 temperatures above current levels are more likely to benefit the latter species. However, the  
65 relationship between temperature and population dynamics for *G. pallida* has not been  
66 examined under different *in vivo* field conditions in the UK. The aim of the present study was  
67 to investigate the multiplication and development of PCN at two field sites in the UK and  
68 compare this with multiplication and development in controlled environments.

69

## 70 Materials and methods

### 71 **Nematodes and plant material**

72 *Globodera pallida* population (Lindley) (Pa2/3) cysts from The James Hutton Institute (JHI)  
73 PCN collection were used for experiments in temperature-controlled conditions. This  
74 population had been maintained in glasshouse conditions (20 °C 16 h, 16 °C, 8 h) for over 30  
75 years at the JHI. Cysts were collected on a 250 µm sieve, randomly selected and packed into  
76 nylon bags containing 30 cysts per bag. Cysts of *G. pallida* (Lindley) were also used for the  
77 validation of qPCR assays.

78 The susceptible cultivar Desirée and the moderately resistant cultivar Morag, which  
79 has resistance to *G. pallida* derived from *Solanum vernei* (Phillips & Trudgill, 1998), were  
80 used in a pot experiment to examine the life cycle of *G. pallida* in temperature-controlled  
81 conditions. For the field experiment, four susceptible cultivars were used at each site  
82 (Desirée, Maris Piper, Cara and Estima) whilst the susceptible cultivar Edzell Blue, which  
83 produces blue tubers, was used to produce guard rows as an additional measure to distinguish  
84 the boundaries of the plots. The four cultivars selected are commonly grown in the UK and  
85 are classified as early (Estima), main crop (Desirée and Maris Piper) and late cropping  
86 (Cara).

87

### 88 **Occurrence of juveniles, males and cysts in the soil under different temperature regimes** 89 **in controlled environments**

90 Growth cabinets were used to monitor the occurrence of hatched juveniles, free-living adult  
91 males and new cysts of *G. pallida* in the soil at two different temperature regimes. The  
92 growth cabinets (Phytotron model 1700; Sanyo) were set as follows: (i) 16 °C for 16 h with

93 light ( $525 \mu\text{mol m}^{-2} \text{s}^{-1}$ ) and 11 °C for 8 h with no light and (ii) 19 °C for 16 h with light and  
94 14 °C for 8 h with no light. Soil temperatures were recorded every 3 h in the pots in the  
95 growth cabinet experiment with DS1920-F5 Temperature iButtons (HomeChip Ltd) placed  
96 into the soil at a depth of 10 cm. The average soil temperatures of these regimes were 14.3  
97 and 17.3 °C, respectively. The relative humidity inside the growth cabinets was maintained at  
98 75%. The accumulated day degrees (DD) were calculated using the formula  $DD = (T_{\text{mean}} -$   
99  $T_{\text{base}}) \times \text{Days}$ , where  $T_{\text{mean}}$  is the mean temperature of that day and  $T_{\text{base}}$  is the base  
100 temperature needed for *G. pallida* to hatch, which was 4 °C according to Ebrahimi *et al.*  
101 (2014).

102 Tuber pieces (3 cm diameter) with a single sprout were cut with a melon scoop from  
103 tubers of cvs Desirée or Morag and planted in pots containing 500 g autoclaved sand:loam  
104 (50:50). Plants (96 per cultivar) were grown in the greenhouse until they were approximately  
105 10 cm high before being moved to the growth cabinets. The plants were arranged in a  
106 randomized design in two growth cabinets for each temperature regime and, 2 days later, a  
107 nylon cyst bag containing 30 cysts was planted into the soil of each pot, at a depth of about 5  
108 cm, beside the tuber piece. Five weeks after inoculation, the cyst bags were removed from the  
109 pots to distinguish between juvenile nematodes originating from these cysts and those from  
110 new females. The bags were removed with tweezers by pinching the top of the bag at the soil  
111 surface and pulling from the soil.

112 Three pots per temperature and cultivar were randomly selected at 7-day intervals for  
113 16 weeks. The juveniles and males were extracted from the soil using the Baermann funnel  
114 method (Viglierchio & Schmitt, 1983). After 2 days, nematodes were collected and the  
115 numbers of juveniles and males were counted with a microscope. Additionally, the total  
116 number of cysts per pot was determined in weeks 8 to 16 by using acetone flotation to

117 recover the cysts that were collected in the sieves used with each Baermann funnel (Brodie *et*  
118 *al.*, 1976).

119

## 120 **Field trials**

121 In 2011, field trials took place in two locations that were naturally infested by *G. pallida*:  
122 East Lothian (lat 56.0170, long -2.8375) and a site near Harper Adams University, Newport  
123 (lat 52.7797, long -2.4275). The soil temperatures were monitored with DS1920-F5  
124 temperature monitoring ibuttons that were placed in the centre of two neighbouring potato  
125 ridges at a depth of 20 cm. Temperatures were recorded every 3 h during the trials.

126         The treatment of 10% oxamyl nematicide granules (55 kg ha<sup>-1</sup>) was applied at the  
127 time of planting potatoes in East Lothian and fosthiazate (30 kg ha<sup>-1</sup>) was applied  
128 immediately before planting at the site near Harper Adams. The farm managers were  
129 responsible for all agricultural operations, including nematicide application, fertilization and  
130 application of other crop protection products at each site.

131         Each site had six experimental blocks and, within each, there were five replicates of  
132 four cultivars (Desirée, Maris Piper, Cara and Estima) in a randomized design. One replicate  
133 consisted of three tubers of each cultivar, planted 25 cm apart. Between trials and at the end  
134 of each row, two tubers of the guard plants (Edzell Blue) were planted 25 cm from the  
135 experimental blocks and from each other. The experiments were set up on 4 April 2011 and 4  
136 May 2011 at the Harper Adams and East Lothian sites, respectively.

137         To estimate the initial populations of PCN, 500 g of soil was taken from each  
138 experimental block with a 15 mL corer. Soil samples were subsequently collected at  
139 approximately monthly intervals, from May to September at Harper Adams and from June to

140 October in East Lothian. During harvesting, approximately 5 L of soil, containing and  
141 surrounding the roots of each of the three plants per replicate, was collected with a spade,  
142 combined and mixed in a bucket. Subsequently, a subsample (at least 500 g) was placed in a  
143 labelled bag. Soil samples were air dried in the glasshouse by spreading each in a plastic tray  
144 (38 × 24 × 5 cm) and, when dry, a 400 g soil sample was packaged in a labelled paper bag  
145 and transferred to Science and Advice for Scottish Agriculture (SASA, UK) for cyst  
146 extraction using their automated soil washing carousel (Meku). The wet filter papers on  
147 which the cysts and debris were collected following their extraction from the soil with the  
148 carousel, were dried and the cysts were further purified by acetone flotation (Brodie *et al.*,  
149 1976) and then transferred into a 2 mL Eppendorf tube<sup>384</sup>

150

151 To monitor nematode development, root samples of cvs Desirée and Cara were  
152 collected when each trial was harvested. The root systems for each replicate of three plants  
153 were cut off and the remaining soil was gently shaken off. Roots were chopped into 5–7 cm  
154 segments and stored in FAA (formalin–acetic–alcohol, 2:1:10) solution (Hooper, 1970).  
155 Later, a  $0.5 \pm 0.2$  g subsample was stained with acid fuchsin (Bridge *et al.*, 1981) and then  
156 examined by microscope (Olympus S7-ST).

157

### 158 **DNA extraction from soil samples**

159 The floats obtained following acetone purification were placed in 2 mL Eppendorf tubes with  
160 two metal beads per tube and pulverized in a mixer mill MM300 (Retsch) for 1.5 min at 30  
161 Hz. The resulting powder was mixed with 0.5 mL GeneScan lysis buffer (Neogen Europe  
162 Ltd) and ground again for 30 s. Samples were centrifuged for 15 s at 15 600 g and then 5  $\mu$ L



163 of 20 mg mL<sup>-1</sup> proteinase K in 40% (v/v) glycerol (Sigma-Aldrich) was added and incubated  
164 for 1 h at 65 °C. After incubation, 0.5 mL chloroform:isoamyl alcohol (24:1) (Sigma-  
165 Aldrich) was added and mixed by inverting the tube five times. Samples were centrifuged for  
166 10 min at 15 600 g in an Eppendorf centrifuge, the upper aqueous phase (450 µL) was  
167 transferred into a new tube and 360 µL of ice-cold propan-2-ol (VWR) was added and mixed  
168 thoroughly. Subsequently, samples were incubated for 30 min at 20 °C, centrifuged for 10  
169 min at 15 600 g and the pellet was retained. The pellet was washed twice with 0.5 mL 75%  
170 ethanol and centrifugation for 5 min at 15 600 g and then resuspended in 100 µL sterile  
171 dH<sub>2</sub>O.

172 The resulting DNA was further purified on PVPP columns. First, a 600 µL suspension  
173 of 10% PVPP (Sigma-Aldrich) was made up with sterile distilled water and transferred to a  
174 spin column (NBS Biologicals) in a 2 mL microcentrifuge tube. Tubes were centrifuged at  
175 11 000 g for 1 min, the catch-tube was emptied and the centrifugation was repeated (the spin  
176 column was rotated 180° within the centrifuge before the second spin). The resuspended  
177 DNA was transferred to the spin column and centrifuged at 11 000 g for 1 min. Purified  
178 eluate was transferred to a new sterile 1.5 mL Eppendorf tube and stored at -20 °C.

179

## 180 **PCN quantification by qPCR**

181 The oligonucleotide design and optimization of specificity and sensitivity of the PCN qPCR  
182 assay was previously described by Reid *et al.* (2010). The primers and probes used were  
183 designed for the rDNA ITS1 region of *G. pallida* and *G. rostochiensis*; primer 1 (forward) 5'-  
184 CGTTTGTGTTGACGGACAYA-3', primer 2 (reverse) 5'-  
185 GGCGCTGTCCRTACATTGTTG-3', *G. pallida* MGB probe 5'-6FAM-  
186 CCGCTATGTTTGGGC-3', *G. rostochiensis* MGB probe 5'-6FAM-

187 CCGCTGTGTATKGGC-3'. All DNA from the field samples was tested for both PCN  
188 species; however, no *G. rostochiensis* was detected when compared to the DNA standard  
189 prepared from *G. rostochiensis* cysts from the JHI collection. To determine whether the assay  
190 was valid over a wide range of cyst densities, DNA was extracted from *G. pallida* (Lindley)  
191 cysts that had been recovered from a 1:1 sand:loam mixture (260 g per pot) with 12 to 644  
192 cysts per pot (average number of eggs  $205.4 \pm 20.3$  per cyst). DNA was extracted from these  
193 cysts as described for the field samples.

194 Real-time qPCR reactions were set up using a Genesis Workstation 150 (Tecan Inc.)  
195 in 96-well plates (Applied Biosystems/Thermo Fisher Scientific) at SASA, UK. The 30  $\mu\text{L}$   
196 reactions contained 15  $\mu\text{L}$  Environmental BLUE Ready Mix (Eurogentec Ltd), 1.25  $\mu\text{L}$  each  
197 of the forward and reverse primers for either species of PCN ( $5 \text{ pmol } \mu\text{L}^{-1}$ ), 1.25  $\mu\text{L}$  of either  
198 the *G. pallida*- or *G. rostochiensis*-specific probe ( $5 \text{ pmol } \mu\text{L}^{-1}$ ), 6.25  $\mu\text{L}$  distilled water  
199 (Sigma-Aldrich) and either 5  $\mu\text{L}$  DNA that had been extracted from samples and diluted 1:10  
200 with  $\text{H}_2\text{O}$ , or 5  $\mu\text{L}$   $\text{H}_2\text{O}$  as a negative control. The reactions were then aliquoted in triplicate  
201 into 384-well plates (Applied Biosystems) using a Genesis robot (Tecan). Standards (in  
202 triplicate) were created from *G. pallida* cyst DNA dilutions of 10, 1, 0.1, 0.01 and 0.001 ng  
203  $\mu\text{L}^{-1}$ . Amplification was performed in an ABI 7900HT (Applied Biosystems) real-time  
204 machine run in the standard mode with the following cycling conditions: 50 °C for 2 min, 95  
205 °C for 5 min, followed by 40 cycles of 95 °C for 15 s and 60 °C for 1 min. Linear regression  
206 of the quantity of copies (qty values) versus egg number and calculation of the corresponding  
207  $R^2$  value were performed using EXCEL v. 16.0.4456.1003 (Microsoft) and GENSTAT v.  
208 16.1.010916.

209 The average value of qty values obtained from the qPCR performed on DNA  
210 extracted from cysts was multiplied by 10 (for the dilution) and fitted to the validated linear  
211 regression model. In order to establish the number of eggs per g soil, the number of eggs

212 obtained from qPCR was divided by 400 g (dry weight of the soil sample from the field). For  
213 standardizing the egg numbers between plots and fields, the data is presented as a  
214 multiplication ratio of the population at the sampling time point divided by the initial number  
215 of eggs in that experimental plot.

216

### 217 **Hatching of field cysts**

218 Cysts were recovered from soil samples collected at the final harvest and subjected to a  
219 period of at least 3 months at 4 °C. Hatching experiments were performed on a thermal  
220 gradient table following the protocol of Kaczmarek *et al.* (2014a, b) with a temperature range  
221 from 5 to 19 °C.

222

### 223 **Statistical analysis**

224 Data was transformed and analysed using GENSTAT v. 17.1 and EXCEL v. 14.0.4760.1000 and  
225 v. 16.0.4456.1003. Data (numbers of J2s, males, cysts and eggs) were analysed using  
226 standard analyses of variance, linear and logarithmic regression as well as *t*-tests as detailed  
227 in the text. The data were subjected to logarithmic transformations where specified. The  
228 hatching data was analysed following the procedures in Kaczmarek *et al.* (2014a, b).

229

## 230 **Results**

### 231 **Occurrence of juveniles, males, and cysts in the soil under two different temperature** 232 **regimes**

233 The first presence of juveniles in the soil was recorded at week 1 for both cvs Desirée and  
234 Morag and in both the 14.3 and 17.3 °C temperature regimes, after 62 and 80 DD  
235 respectively. This was followed by an increase in numbers and then a subsequent decrease  
236 after week 2 (between 104 and 295 DD; Fig. 1a,b). Around 1219 DD (week 10 at 17.3 °C),  
237 increasing numbers of J2s were recorded with Desirée (Fig. 1b) until the end of the  
238 experiment at week 16 (1500 DD), with comparable numbers ( $51 \pm 23$  J2 per pot) to those  
239 observed during the first hatching peak. A small and later second hatch was also observed at  
240 14.3 °C at 1019 DD (week 14; Fig. 1a) with Desirée. Juveniles recovered from the pots with  
241 Morag, kept the same hatching trend as susceptible Desirée. The fewest numbers of juveniles  
242 were observed for Morag at the cooler temperatures.

243 The first appearance of males in both temperature regimes was at week 5 (332 DD)  
244 (Fig. 1c, d) with the highest numbers of males at 14.3 °C on Desirée at week 6 (436 DD)  
245 whereas fewer were observed on Morag at 14.3 °C at week 6. At 17.3 °C, males were  
246 observed again at weeks 11, 13, 14, 15 and 16 with Desirée and at week 15 with Morag.  
247 Males were not observed at 14.3 °C for either cultivar after week 11.

248 Mature (tanned) cysts were recovered from the soil from 9 weeks after inoculation  
249 onwards with Desirée at 14.3 °C (655 DD) and 17.3 °C (844 DD), with greater numbers at  
250 the higher temperature (Fig. 1e), peaking at 938 DD. Very few cysts were recovered with  
251 Morag at either temperature. There was a significant difference in the overall mean number  
252 of cysts recorded between the two temperature regimes (14.3 and 17.3 °C), the mean  
253 responses at different temperatures ( $P = 0.005$ ) and for different cultivars ( $P < 0.001$ ). There  
254 was also an interaction between temperature and cultivar ( $P = 0.007$ ).

255

256 **Field trials**

257 *Soil temperatures*

258 The average soil temperatures at a depth of 20 cm in the potato drills over the growing  
259 seasons of 147 and 157 days were 14.1 and 15.5 °C at the East Lothian and Harper Adams  
260 sites, respectively (Table 1). The fluctuations in the soil temperature during the growing  
261 season at the 2 sites is shown in Fig. 2. The temperatures at Harper Adams were higher than  
262 at East Lothian during three of the periods between soil samplings and were particularly high  
263 during the final harvest period (average 16.6 compared to 12.9 °C). Assuming a base  
264 temperature of 4 °C for *G. pallida* (Ebrahimi *et al.*, 2014), there were 1511 and 1804 DD at  
265 the East Lothian site and Harper Adams sites respectively. A *t*-test confirmed significant  
266 differences in the soil temperature between the sites ( $P < 0.001$ ).

267 *qPCR validation*

268 The primers and probes for the qPCR assay were designed to be specific to *G. pallida* or *G.*  
269 *rostochiensis* (Reid *et al.*, 2010) and were based on rDNA ITS sequences of many PCN  
270 isolates from Europe. All samples were tested for both PCN species but no *G. rostochiensis*  
271 was detected. In order to use this assay to quantify *G. pallida* in the field trial soil samples, it  
272 was necessary to establish that the relationship between egg numbers and qPCR qty was  
273 linear over a wide range of eggs numbers. The DNA yield (qPCR qty) from *G. pallida* cysts  
274 was positively correlated with the number of eggs (Fig. 3) giving the linear regression model:  
275  $y = 25.367x$  with  $R^2 = 0.7619$ , where  $y$  is the number of eggs of *G. pallida* and  $x$  is DNA yield  
276 multiplied by 10 for the dilution. Once established, the model was applied to field samples to  
277 determine the egg number per sample.

278 *Quantifying PCN in the East Lothian field trial*

279 The initial population ( $P_i$ ) of the East Lothian field site was an average of  $11.7 \pm 1.6$  (without  
280 nematicide plots) and  $20.6 \pm 2.2$  (with nematicide plots) eggs per g soil of *G. pallida*.  
281 Changes in the *G. pallida* population over the growing season, without and with nematicide  
282 treatment for the four cultivars are presented in Figure 4a and 4b, respectively. In the plots  
283 without nematicide treatment, the multiplication ratio (population at sampling:initial  
284 population) generally decreased at 4, 8 and 12 weeks, which is consistent with a decrease in  
285 the egg content of cysts due to hatching. At week 16, the multiplication ratio peaked. At week  
286 20 (1511 DD) the final  $P_f/P_i$  (final population/initial population) dropped significantly and  
287 ranged from  $5.0 \pm 1.9$  for Cara to  $2.4 \pm 0.4$  for Desirée. However, in the nematicide-treated  
288 plots, the multiplication remained at similar levels over the whole growing season. The  
289 overall trend for the effect of nematicides is shown in Figure 4c, where the multiplication  
290 ratio at each sampling time has been averaged over the four cultivars. The final average  $P_f/P_i$   
291 values were  $3.6 \pm 0.6$  for the untreated and  $0.7 \pm 0.1$  for nematicide-treated plots. The relative  
292 increase in the population was clear without nematicide treatment at weeks 16 and 20 in  
293 contrast to the nematicide treatment where the final population was lower than the initial  
294 population. To assess the influence of nematicide treatment and cultivar on multiplication  
295 ( $P_f/P_i$ ), an ANOVA was used. It revealed no significant differences in the reproduction ratio  
296 between the cultivars during the growing season; however, there nematicide treatment had a  
297 significant influence on population multiplication ( $P < 0.001$ ) over the growing season.

#### 298 *Quantifying PCN in the Harper Adams field trial*

299 The  $P_i$  of *G. pallida* in the Harper Adams field trials was estimated prior to planting in April.  
300 The average of  $9.9 \pm 0.6$  (without nematicide) and  $6.3 \pm 0.9$  (with nematicide) eggs per g soil  
301 was determined by qPCR. Samples were negative when tested for *G. rostochiensis*. Changes  
302 in the *G. pallida* population over the growing season in the plots without and with nematicide  
303 treatment for the four cultivars are presented in Figure 5a and 5b respectively. For samples

304 without nematicide treatment, there were generally slight reductions in the population at 4  
305 and 8 weeks after planting compared to preplant levels. At week 16 (1193 DD), the  
306 multiplication ratio increased to reach the Pf/Pi at final harvest of  $7.7 \pm 3.0$  for Cara and  $2.5 \pm$   
307  $0.5$  for Estima. For the nematicide treatment, the levels remained similar for the first 12  
308 weeks of the field experiment, but the multiplication ratio increased at the fourth sampling  
309 time and at the combined harvests at weeks 20 and 22. The final average Pf/Pi values for the  
310 four cultivars were  $5.2 \pm 1.0$  and  $2.1 \pm 0.3$  for the untreated and treated plots, respectively  
311 (Fig. 5c). To assess the influence of nematicides and different cultivars on multiplication, an  
312 ANOVA was performed. There were no significant differences in the Pf/Pi between the  
313 cultivars during the growing season; however, there was a significant influence of nematicide  
314 treatment on Pf/Pi ( $P < 0.001$ ) over the growing season.

#### 315 *Relationship between initial population and reproductive factor (Pf/Pi)*

316 The Harper Adams site was lightly to moderately infested with *G. pallida* (<20 eggs per g  
317 soil) and the site in East Lothian varied from lightly to heavily infested with a range of initial  
318 population densities (5–50 eggs per g soil). The results indicated that in the untreated plots at  
319 both sites, particularly at East Lothian, there was a trend towards a decreasing multiplication  
320 rate with increasing initial population (Fig. 6a). The multiplication rate was significantly  
321 higher in the plots with lower initial densities ( $P < 0.001$ ). Also, the multiplication rate (Pf/Pi)  
322 was reduced with the oxamyl treatment at the East Lothian site. At the Harper Adams trial, no  
323 trend was found between Pi and Pf/Pi in the nematicide treated plots (Fig. 6b). Further  
324 statistical tests (ANOVA) on the multiplication rate revealed a significant effect of initial  
325 population on the Pf/Pi ( $P < 0.001$ ) and nematicide treatment ( $P < 0.001$ ) at both sites but no  
326 difference in multiplication rate associated with different cultivars.

#### 327 *Developmental stages of the nematodes in the roots from the field trials*

328 At the first harvest (week 4; 281 DD) from the East Lothian site, J2 nematodes were most  
329 frequently observed in the roots in both nematicide-treated and untreated samples, although  
330 single J3 and J4 stages were also found (Fig. 7a, b). At week 8 (593 DD), mainly J3 stage  
331 nematodes were observed, and much lower numbers of J2 and a few J4 stage nematodes.  
332 There was an increase in the J2 nematodes at week 12 (895 DD), particularly in Desirée  
333 roots.

334 At Harper Adams, J2 nematodes were observed in the roots of both Desirée and Cara  
335 at week 4 (278 DD; Fig. 8a, b). The highest number was recorded in Cara from the untreated  
336 plots. At week 8 (544 DD), both J3 and J4 stage nematodes were observed. J2 and J3 stages  
337 were observed in Cara samples from week 12 (856 DD) in non-nematicide-treated plots,  
338 while no nematodes were found in roots from the treated plots. Juvenile stage 2 nematodes  
339 were observed to increase in numbers in untreated plots at weeks 20 and 22 (end of  
340 experiment 1804 DD).

341 In the East Lothian root samples, the first occurrence of females and highest numbers  
342 were found on roots from Cara and Desirée 8 weeks (593 DD) after planting (Fig. 7c). Lower  
343 numbers of females were seen at week 12 (895 DD); however, at week 16 (1189 DD) no  
344 females were observed. Surprisingly the highest numbers of females were found with the  
345 nematicide treatment. At week 20 the roots had deteriorated, and it was not possible to  
346 examine them histologically. A few females were observed on the roots of Cara at week 4  
347 (278 DD) at the Harper Adams site, but most were seen at 8 (544 DD) and 12 weeks (856  
348 DD), with higher numbers observed without nematicide treatment (Fig. 8c). Females were  
349 found at combined weeks 20 and 22 (1804 DD) on the roots of Cara and Desirée in the  
350 untreated samples, and a few were seen at week 22 (1804 DD) in roots from the nematicide  
351 treatment with both cultivars.



352

### 353 **Hatching test with field cysts**

354 The cumulative proportion of eggs that hatched from the field cysts from the two sites was  
355 calculated for each of the sets of cysts over the 56 days of the experiment (Fig. 9a) and fitted  
356 to a linear regression model. Analysis of variance on the log-transformed curve parameters  
357 suggested no evidence of differences in the means between the two sites (East Lothian and  
358 Harper Adams) for maximum hatch rate or time to 50% hatch. There was a (marginally)  
359 significantly higher total proportion hatching at East Lothian than Harper Adams ( $P < 0.001$ )  
360 (Fig. 9b). This is confirmed by ANOVA on the final counts (raw data), where the mean total  
361 hatch at East Lothian was 975 and for Harper Adams 711 ( $P = 0.03$ ), and ANOVA on the  
362 total proportion hatching. There are also significant differences in the time to 50% hatch and  
363 total hatch between temperatures but not for hatching rate.

364

### 365 **Discussion**

366 The main goal of these experiments was to investigate the relationship between the life cycle  
367 of *G. pallida* and temperature and to assess the potential for the development of a second  
368 generation of *G. pallida* in UK fields. The data has also been used for the development of a  
369 dynamic stage-structured simulation model (Kettle & Nutter, 2015) and for risk assessments  
370 in relation to climate change (Skelsey *et al.*, 2018). The growth cabinet experiments provide a  
371 temporal framework for the rates of development in different temperature regimes and  
372 indicate the relative abundance of the different life stages in different temperature regimes.  
373 The pot experiment was performed to allow comparison between the life cycle of *G. pallida*  
374 in a controlled environment and in more variable field conditions. There was no effect of the

375 potato cultivar on the number of juveniles recovered from the soil in the initial hatching peak.  
376 This is in agreement with Turner (1990) who found no significant difference in hatching of  
377 *G. pallida* with various *S. vernei* hybrids. In previous *in vitro* hatching experiments  
378 conducted in petri dishes using potato root diffusate to induce hatching (Kaczmarek *et al.*,  
379 2014a, b) , newly hatched juveniles of *G. pallida* were first observed at day 9; however,  
380 more hatched by day 12 at 13–25 °C. The results obtained with the growth cabinet  
381 experiments using potato plants in soil, showed a similar delay before hatching occurred  
382 observed by the presence of juvenile nematodes recovered from the soil, and hatching  
383 continued for several weeks. Adult males were recovered from the soil at 333 DD and the  
384 maximum number of males was recorded about 430 DD in both temperature regimes;  
385 however, more males were recorded for the growth cabinet with the lower temperature.  
386 Further investigation is needed to determine if there is an effect of temperature on sex  
387 determination (Jones *et al.*, 2017; Skelsey *et al.*, 2018).

388         In the present investigation, new cysts were observed in the 17.3 °C regime from nine  
389 weeks after inoculation (844 DD), followed by the start of a second increase in the number of  
390 juveniles in the soil at week 10 (938 DD), indicating that fewer than 10 weeks were required  
391 for one generation to be completed and for another to start in these conditions for this  
392 population of *G. pallida*. The juveniles observed at 10 weeks (938 DD) and later are  
393 consistent with the hatching of a second generation originated from new, first generation  
394 females; this implies that diapause was not obligatory for all juveniles from the first  
395 generation, although the amount and timing of the second hatch was affected by the  
396 temperature regime. Hatching with PCN is not tightly synchronized, resulting in the  
397 occurrence and overlapping of different developmental stages over several weeks.  
398 Nevertheless, based on these results, if the second generation develops at the same rate as the  
399 first generation at this temperature, then second generation cysts would be expected at week

400 18 (about 8 weeks after the second appearance of juveniles in the soil). Because it was not  
401 possible to sustain the plants for longer than 16 weeks in the growth cabinets despite regular  
402 fertilization and watering, it was not possible to determine whether a second increase in cysts  
403 would occur.

404 In both temperature regimes, lower numbers of cysts were observed with partially  
405 resistant Morag compared to Desirée, consistent with previous reports (Phillips *et al.*, 1980;  
406 Phillips & Trudgill, 1998). However, there was a much greater response to temperature with  
407 susceptible Desirée. This indicates that partially resistant cultivars may be more effective  
408 than susceptible cultivars in controlling population multiplication over a wider temperature  
409 range. Significantly lower egg numbers per cyst were found with Morag than with Desirée,  
410 but numbers did not differ significantly between the two temperature regimes. This contrasts  
411 with other reports (Bendezu *et al.*, 1998; Da Cuhna *et al.*, 2012) where resistance from *S.*  
412 *vernei* did not significantly reduce the number of cysts or egg content in more virulent  
413 populations of *G. pallida* from Portugal and Bolivia, although with this source of resistance,  
414 less virulent populations of *G. pallida* had significant differences in their multiplication.

415 The experiments in temperature-controlled environments described here used a PCN  
416 population from the JHI collection that has been multiplied in glasshouse conditions and may  
417 have become adapted to these conditions. Other work has noted differences between  
418 populations of *G. pallida* in their temperature responses (Foot, 1978) and a population of *G.*  
419 *rostochiensis* was selected that matured more quickly at low temperatures following several  
420 years of early cropping in Ayrshire, UK (Ellenby & Smith, 1975). However, the hatching  
421 tests over a range of temperatures performed in the present study with *G. pallida* cysts  
422 isolated from two field sites confirmed the responses obtained previously with glasshouse  
423 cultured *G. pallida* (Kaczmarek *et al.*, 2014 a, b).

424           The qPCR assay for *G. pallida* and *G. rostochiensis* (Reid *et al.*, 2010) was used to  
425 monitor the densities of eggs per g soil in soil samples taken at the two field sites. This is the  
426 first time this assay has been used to monitor PCN population dynamics in the field over a  
427 growing season and demonstrates the value of this high-throughput assay for assessing the  
428 impact of different PCN control programmes. The relationship between egg numbers and  
429 DNA yield was validated using *G. pallida* cysts recovered and quantified from soil using the  
430 same extraction procedure used for the field samples.

431           The mean soil temperature between planting and first harvest was 13.9 °C at the  
432 Scottish site compared to 14.4 °C at the English site, below the optimal temperatures that  
433 have been observed for hatching (Kaczmarek *et al.*, 2014a, b). At harvest 1 at East Lothian  
434 but not in the Harper Adams root samples, J3 stage nematodes were observed and a few  
435 female nematodes were observed on the roots from Harper Adams. The mean soil  
436 temperatures in sampling periods 1 and 2 were lower at Harper Adams than the East Lothian  
437 site; however, they increased to 15.5 and 16.6 °C in sampling periods 4 and 5 giving 1193  
438 and 1804 DD respectively, whereas they dropped to 14.4 and 14.1 °C at the East Lothian site  
439 with 1189 and 1511 DD respectively. Females were observed on the roots from week 4  
440 onwards at Harper Adams with relatively warmer soil temperatures (14.8–15.5 °C) as found  
441 by Jones *et al.* (2017). It is probable that a second generation developed, as both juveniles  
442 and females were observed at harvests 5 (week 20) and 6 (22 weeks), although the possibility  
443 that the late females observed in the field samples may have resulted from a delayed initial  
444 hatch cannot be discounted. However, two generations of *G. pallida* in a growing season has  
445 been reported with high numbers of cysts in soil temperatures from 18 to 22 °C in Avezzano,  
446 Italy (Greco *et al.*, 1988) and in the present study, a second hatch was observed at 1219 DD  
447 in the controlled environment cabinet at 17.3 °C. Ebrahimi *et al.* (2014) estimated the degree  
448 day requirement for *G. pallida* of 450 DD with a 4 °C base temperature. There were 1511 DD

449 and 1842 DD at the East Lothian and Harper Adams sites respectively, both sufficient for two  
450 generations to be completed.

451           The general trend of increasing temperatures associated with climate change  
452 combined with the warmer conditions that occur in some regions in the UK are making it  
453 more likely that conditions that support two generations of *G. pallida* within a growing  
454 season will occur. A recent report showed that both species of PCN are developing more  
455 quickly in Belgium than in the past (Ebrahimi *et al.*, 2014) and this has resulted in a change  
456 of advice for the time of harvesting early cultivars to prevent PCN from completing its life  
457 cycle. In 2014 in Lincolnshire, average soil temperatures at a depth of 20 cm of 15.3 °C were  
458 recorded and many females were observed developing on roots of susceptible untreated  
459 potatoes 21 weeks after planting (A. Barker, Barworth Agriculture LTD, Sleaford, personal  
460 communication). It is not known whether these late developing females matured into cysts or  
461 whether they contained viable eggs that could contribute to a higher final Pf/Pi; this requires  
462 further investigation. In some conditions, a partial second generation, where a substantial  
463 number of nematodes hatch but are not able to complete the second generation of mature  
464 cysts, could lead to lower final Pf/Pi values and a trap crop effect. An effect of temperature on  
465 in-egg mortality and spontaneous hatching, affecting population dynamics, has also been  
466 reported (Devine *et al.*, 1999).

467           Nematicides can affect PCN during different developmental stages, mainly affecting  
468 juvenile nematodes before they reach the potato plants. Fosthiazate temporarily inhibited  
469 hatching of *G. pallida* in an *in vitro* test and suppressed hatching in soil (Woods *et al.*, 1999).  
470 The results of the nematicide trial in the current investigation are consistent with other studies

471           that have reported that carbamate nematicides do not permanently stop hatching in  
472 field conditions (Evans & Wright, 1982; Woods *et al.*, 1999); nevertheless, they do delay the

473 process significantly. Although the decline in egg numbers was less in nematicide treated  
474 plots than in untreated plots, this was probably due to less hatching with the nematicide  
475 treatment. Generally, fewer J2 stage nematodes were observed in the roots of Cara and  
476 Desirée with nematicide treatment, consistent with a report by Minnis *et al.* (2004) and fewer  
477 females were observed at week 8 with the nematicide treatments at both sites. At the East  
478 Lothian site there was significantly lower multiplication with the oxamyl treatment ( $P <$   
479  $0.001$ ). This is in contrast to the results of Minnis *et al.* (2004) that showed no significant  
480 difference in PCN multiplication between fields treated or untreated with oxamyl. However,  
481 as mentioned previously, in the current investigation, the density of the initial population in  
482 untreated plots ranged from 5 to 20 eggs per g soil while the plots for nematicide treatment  
483 had  $>20$  eggs per g of soil; therefore, it is difficult to differentiate between a reduction in the  
484 multiplication rate due to the nematicide treatment or due to the higher initial number of eggs  
485 in the soil.

486         Understanding how climate change may drive the life cycle and population dynamics  
487 of PCN is important because of the role of these plant pests in potato yield reduction; in  
488 addition, the management of this pest is of increasing concern due to the withdrawal of  
489 nematicides from use. The results indicate that the development of multiple generations of  
490 PCN within one growing season in the UK is more likely with increasing soil temperatures  
491 associated with climate change and this could have significant effects on management of  
492 population multiplication and prevention of the spread of PCN (Jones *et al.*, 2017; Skelsey *et*  
493 *al.*, 2018).

494

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507

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595

## 596 **Figure legends**

597 **Figure 1** Comparison of number of *Globodera pallida* juveniles at 14.3 °C (a) and 17.3 °C  
598 (b), males at 14.3 °C (c) and 17.3 °C (d), and cysts at 14.3 and 17.3 °C (e) recovered from the  
599 soil over 16 weeks in the growth cabinet experiment with susceptible cv. Desirée and  
600 partially resistant cv. Morag. The number of degree days (DD) at each sampling time are  
601 given in brackets. Two pots were used for each temperature and cultivar combination, and  
602 nematodes were extracted with a Baermann funnel. The bars indicate the standard error of  
603 means.

604 **Figure 2** Soil temperatures at 20 cm in potato drills over the growing season in Harper  
605 Adams (4 April 2011 to 6 September 2011, 155 days) and East Lothian (3 May 2011 to 28  
606 September 2011, 147 days). Soil temperatures were taken every 3 h in two adjacent drills and  
607 the averages are shown.

608 **Figure 3** Linear regression between egg numbers of *Globodera pallida* and quantity of  
609 copies determined by qPCR. The qPCR results for 11 samples were averaged across three  
610 qPCR replicates.

611 **Figure 4** Changes in the multiplication ratio (population at sampling time/initial population)  
612 of *Globodera pallida* over the growing season (a) without nematicide treatment, (b) with  
613 nematicide treatment, and (c) average multiplication ratio for the four cultivars (Cara,  
614 Desirée, Estima and Maris Piper) with and without nematicide treatments at the East Lothian  
615 site. Soil samples were taken at planting and at 4-weekly intervals (degree days (DD) are  
616 shown in brackets after each sampling time) and the eggs per g soil determined from isolated  
617 cysts using qPCR. The bars are standard errors of the means for each cultivar.

618 **Figure 5** Changes in the multiplication ratio (population at sampling time/initial population)  
619 of *Globodera pallida* over the growing season (a) without nematicide treatment, (b) with  
620 nematicide treatment and (c) average final population/initial population (Pf/Pi) for the four  
621 cultivars (Cara, Desirée, Estima and Maris Piper) with and without nematicide treatments at  
622 the Harper Adams site. Soil samples were taken at planting and at 4-weekly intervals (degree  
623 days (DD) are shown in brackets after each sampling time) and the eggs per g soil determined  
624 from isolated cysts using qPCR. The bars are standard errors of the means for each cultivar.

625 **Figure 6** Reproductive factor (ratio of final population/initial population (Pf/Pi)) versus  
626 initial population (Pi) of *Globodera pallida* at (a) East Lothian and (b) Harper Adams sites.

627 **Figure 7** The numbers of J2, J3, J4 and females of *Globodera pallida* observed inside 100 g  
628 of acid fuchsin-stained roots of cultivars Cara and Desirée from the 2011 field trials at East  
629 Lothian. (a) Non nematicide-treated plots, (b) nematicide-treated plots, and (c) females from  
630 both treatments. Root samples were examined at harvests 1, 2, 3 and 4 (weeks 4, 8, 12 and  
631 16, respectively, which correspond to 281, 593, 895 and 1189 degree days (DD)). The bars  
632 indicate the standard error of the mean.

633 **Figure 8** The numbers of J2, J3, J4 and females of *Globodera pallida* observed inside 100 g  
634 of acid fuchsin-stained roots of cultivars Cara and Desirée from 2011 field trials at Harper  
635 Adams; (a) non-nematicide-treated plots, (b) nematicide-treated plots, (c) females from both  
636 treatments. Root samples were examined at harvests 1, 2, 3, 4, 5 and 6 (weeks 4, 8, 12, 16, 20  
637 and 22, respectively, which correspond to 278, 544, 856, 1193 and 1804 degree days). The  
638 bars indicate the standard error of the mean.

639 **Figure 9** Proportion of hatched eggs of *Globodera pallida* from the field populations in  
640 potato root diffusate, during 55 days incubation at different temperatures; (a) cumulative and  
641 (b) percentage of total hatch. Bars indicate standard errors of means.