

1 **Short Research Article**
2 **Pollen viability and fruit yield of coriander**
3 **genotypes inoculated with different**
4 **concentrations of *Meloidogyne incognita* race 1**

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6
7
8 **ABSTRACT**
9

Coriander is among the most consumed leafy vegetables in Brazil, employing a large number of people in its production chain. Among the limiting factors to the production of this horticulture there is the *Meloidogyne* caused by gall nematodes. The present work was carried out aiming to understand the impact on coriander plants submitted to the parasitism of *M. incognita* race 1. The number of galls were verified at 30 days after inoculation as well as the plant survival up to the reproductive stage. In addition, pollen viability and fruit yield in coriander plants of two cultivars (Verdão and HTV Dom Luiz), inoculated at sowing with six inoculum concentrations (0, 1,000, 2,000, 4,000, 8,000 and 16,000 eggs / cell) and evaluated in a randomized complete block design with four replications were also analyzed. The plot was composed of one plant. The presence of the pathogen did not influence the pollen viability by means of the acetic Carmine and Alexander dyes. However, neither tetrazolium nor in vitro pollen germination means were efficient in the viability identification. Concentrations of 8,000 and 16,000 eggs/cell did not allow the development of plants, leading them to death. Inoculation at sowing, and evaluation of the number of galls at 30 days did not limit the reestablishment of the plant development and fruit yield, up to the concentration of 4,000 eggs/cell.

Comment [EMF1]: Horticultural crop

10
11 *Keywords: Coriandrum sativum L., inoculum levels, Meloidogyne, Dyestuff.*
12

13 **1. INTRODUCTION**
14

15 Gall nematodes (*Meloidogyne* spp.) are widely distributed phytopathogens in tropical,
16 subtropical and temperate regions of the planet. These worms are classified as one of the
17 five main groups of economically destructive pathogens that affect world food production [1].
18 Within the phytonematological specialty, the *Meloidogyne* species constitute the most
19 economically relevant group, due to the high degree of polyphagy and wide geographic
20 distribution, being constant threats to the rural producers. The species with the most
21 prominence within the genus for presenting themselves as the most cosmopolitan,
22 polyphagous, and weedy to agriculture are: *M. arenaria*, *M. hapla*, *M. incognita* and *M.*
23 *javanica*. The last two are the most important from an economic point of view [2].
24

25 The species belonging to the genus *Meloidogyne* are obligatory endoparasites and the
26 second stage juveniles (J2) are infectants, which penetrate the root elongation zone and
27 migrate intercellularly until they reach the differentiating vascular zone, where they choose
28 from 5 to 7 cells and insert their stylus injecting secretions to induce hypertrophy and
29 hyperplasia forming the galls and establishing the feeding site. From that stage on, the
30 pathogen starts to extract nutrients and photoassimilates from nearby tissues, mainly xylem
31 and phloem [3]. Consequently, they promote, besides gall formation, leaf yellowing,
32 defoliation, retarded growth and wilting as a symptom, which collectively reduce plant vigor
33 and cause mass loss and quality. It is estimated that 12.3% of the annual losses in

34 agricultural production is due to the attack of gall nematodes, causing an economic loss of
35 around US \$ 157 billion [1].

36
37 In addition to underdevelopment, the formation of coenocytes, induced by gall nematodes,
38 acts as a metabolic drain resulting in physiological imbalances, that is, a lack of macro
39 and/or micronutrients [2]. Boron is a micronutrient that plays an important role in the growth
40 of the pollen tube [4], and the lack of this element is a factor that can reduce pollen viability.
41 In this context, plants parasitized with gall nematodes, due to the presence of physiological
42 disturbances, may undergo influence on the viability of the pollen grains resulting in reduced
43 productivity. To verify pollen viability, various techniques can be adopted, which can be
44 grouped in methods of direct in vitro determination [5]; direct in vivo determination [6]; and
45 indirect methods such as staining from the biochemical reaction [7].

46
47 Coriander is among the various cultivated species attacked by gall nematodes. There is no
48 cultivar in Brazil that is indicated as resistant to such pathogens, being necessary the search
49 for superior genotypes. According to [8] the development of breeding programs aimed at
50 resistance to phytonematodes is a process that requires adequate resources and
51 methodologies. Currently, evaluation of the reaction of coriander genotypes to gall
52 nematodes consists of sowing, inoculation at 15 days after sowing and evaluation at 45 days
53 of inoculation, by destructive method for the extraction of pathogen eggs aiming at
54 estimating the reproduction factor (RF) [9, 10, 8, 11].

55
56 The use of a methodology that does not promote the destruction of the plant, allowing the
57 selection among and within segregating populations in the same selective cycle, may
58 promote the attainment of promising results. Thus, the present work aims to verify the
59 development of plants of two coriander cultivars inoculated at sowing with different
60 concentrations of *M. incognita* race 1 inoculum and transplanted after 30 days, evaluating
61 plant survival, percentage of viable and germinated in vitro pollen, and fruit yield.

62 63 64 **2. MATERIALS AND METHODS**

65
66 The experiment was conducted under greenhouse conditions, in the Department of
67 Agronomy of the Federal Rural University of Pernambuco (UFRPE), located at 8°54'47"S,
68 34°54'47"W, 6m high, from July to October 2017. The monthly average temperatures
69 recorded by the weather station of Recife Curado (automatic) varied in average between
70 22.9 - 28.6°C, for the minimum and maximum temperature, respectively [12]. The design
71 was of randomized blocks in a factorial scheme of 2 (cultivars) X 6 (inoculum concentrations)
72 with four replications, whose plot was composed by one plant.

73
74 The reactions of the cultivars Verdão and HTV Dom Luiz were evaluated and handled in six
75 concentrations (0, 1,000, 2,000, 4,000, 8,000 and 16,000) of *M. incognita* race 1. Sowing
76 was carried out in expanded 128-cell polystyrene trays containing commercial substrate. The
77 trays were irrigated, followed by inoculation according to each treatment. Each block
78 contained a plot with Santa Clara tomato cultivar (*Solanum lycopersicum* L.), with a
79 susceptibility standard to gall nematodes, in order to verify the efficiency of the inoculum
80 used.

81
82 The inoculum was obtained from sources kept in tomatoes, Santa Clara cultivar, using the
83 [13] and modified by [14] for the extraction of eggs of *M. incognita* race 1.

84

85 Irrigation was performed according to the water requirement of the crop, without drainage to
86 prevent egg leaching. After germination, fertigation with nutrient solution containing macro
87 and micronutrient was carried out three times a week, adapted from the proposal by [15],
88 taking the same care not to drain and prevent consequent inoculum loss.

89
90 After germination, thinning was carried out leaving only one plant per cell. After 30 days of
91 sowing and inoculation, the roots of each plant of the plot were carefully washed in still
92 water, to remove the substrate without damaging the root system, quantifying the number of
93 galls in the root system and transplanting the seedlings (Figure 1) into 2L pots, properly
94 identified (cultivar and inoculated concentration), containing a substrate based on soil and
95 humus mixture in a ratio of 3:1. These procedures were carried out at around 4:30 p.m., in
96 order to favor seedling adaptation. Irrigation was carried out shortly after transplantation.

97
98 In the pots, the plants were irrigated daily according to the water requirement, and fertigation
99 was applied three times a week. The survival of the plants was observed until the fruits were
100 harvested.

101
102 At the beginning of flowering, when all the plants were flowering, flowers were collected to
103 quantify the viable pollens in each cultivar as a result of the concentration of the pathogen,
104 seeking to verify if the presence and quantity of the pathogen influence pollen viability.

105
106 The pollen was stained with acetic carmine [16], Alexander's solution [17] and tetrazolium
107 salt solutions at concentrations of 0.25%; 0.50%; 0.75% and 1.0% [18]. Each dye reacts with
108 a certain compound and/or structure of the pollen grain, promoting or not promoting grain
109 staining. The stained grains are those considered viable. Acetic carmine indicates
110 chromosomal integrity; Alexander solution contains acid fuchsin and green malachite which
111 react with pollen wall protoplasm and cellulose [19], respectively, whereas tetrazolium salt
112 provides an indication of the metabolic activity of the pollen grain , allowing the estimation of
113 its viability [20] through the reaction of salt with hydrogen resulting from cellular respiration
114 with red pollen [21] indicating the presence of functional enzymes such as peroxidase,
115 esterase and dehydrogenase [22].

116
117 The flowers were collected after the anthesis from 7:00 a.m. to 9:00 a.m., stored in identified
118 paper bags, and taken to the floriculture laboratory where the slides were prepared for
119 visualization and counting of viable pollens in result of the dye. For each dye, four slides per
120 treatment (cultivar X inoculum) were prepared. The pollen of one flower was placed with the
121 aid of a brush on the slide, and then two drops of dye were added before placement of the
122 cover slip. The pollen was observed after 10 minutes of dye addition [19] under a 40X
123 magnification microscope.

124
125 To evaluate the tetrazolium solutions, prepared by diluting the salt in distilled water, four
126 flowers of each treatment were collected for each tetrazolium solution. With a pair of
127 tweezers, the stamens of each flower were placed into an eppendorf, duly identified,
128 containing 1 mL of a certain tetrazolium solution (0.25%, 0.50%; 0.75% and 1%). After that,
129 the tubes were agitated for 20 seconds so that the pollens had contact with the solution. The
130 tubes were then wrapped with aluminum foil and kept at 25°C for 24 hours in a BOD
131 incubator [23, 18]. After this period, the solution of each tube was collected individually with
132 a Pasteur pipette, placed onto a slide, and visualized in a 40X magnification microscope.

133
134 Proposed media were used for pollen germination in vitro for eggplant, citrus and some
135 adaptations because no medium was found for coriander or other species of the Apiaceae
136 family. For all media, phytigel was used instead of agar, leaving the medium more
137 translucent, favoring pollen visualization.

138

139

Table 1. Media used aiming to germinate the coriander in vitro pollen grains

Medium	Reagents						Reference
	Ca(NO ₃) ₂	H ₃ BO ₃	Sacarose	Phytigel	MgSO ₄	KNO ₃	
A	800 mg/L	200 mg/L	100 g/L	10 g/L	-	-	Salles et al., 2006
B	500 mg/L	120 mg/L	100 g/L	10 g/L	120 mg/L	100 mg/L	Tatis et al., 2013
C	4 g/L	3 g/L	10 g/L	10 g/L	-	-	-
D	4 g/L	3 g/L	10 g/L	-	-	-	-
E	-	40 g/L	200 g/L	10 g/L	-	-	-
F	-	40 g/L	100 g/L	10 g/L	-	-	-
G	20 g/L	40 g/L	200 g/L	10 g/L	-	-	-
H	20 g/L	40 g/L	200 g/L	-	-	-	-
I	20 g/L	40 g/L	400 g/L	-	-	-	-
J	20 g/L	20 g/L	400 g/L	10 g/L	-	-	-
L	10 g/L	5 g/L	50 g/L	10 g/L	-	-	-
M	10 g/L	5 g/L	10 g/L	10 g/L	-	-	-
N	5 g/L	5 g/L	10 g/L	10 g/L	-	-	-

140 For both dyes and in vivo media, 150 pollens per slide were evaluated, constituting the experimental
 141 unit.

142

143 After flowering and fruit filling phase, when they were dry, all fruits of each plant were
 144 harvested individually and stored in identified paper bags. Then, the number of fruits of each
 145 experimental plot was quantified, and 100 fruits of each plot were weighed on a precision
 146 scale.

147

148 The number of galls in the root system, number of fruits and weight of 100 fruits variables
 149 were transformed by \sqrt{x} to meet the assumptions of the analysis of variance, and then
 150 submitted to ANOVA and Scott-Knott grouping test at 5% probability. Regression analyzes
 151 were performed for the decompositions within inoculum concentrations. The analyses were
 152 performed using the statistical software SISVAR [24], and the regression graphs were
 153 significant at 1% or 5%, elaborated in Excel.

154

155

156

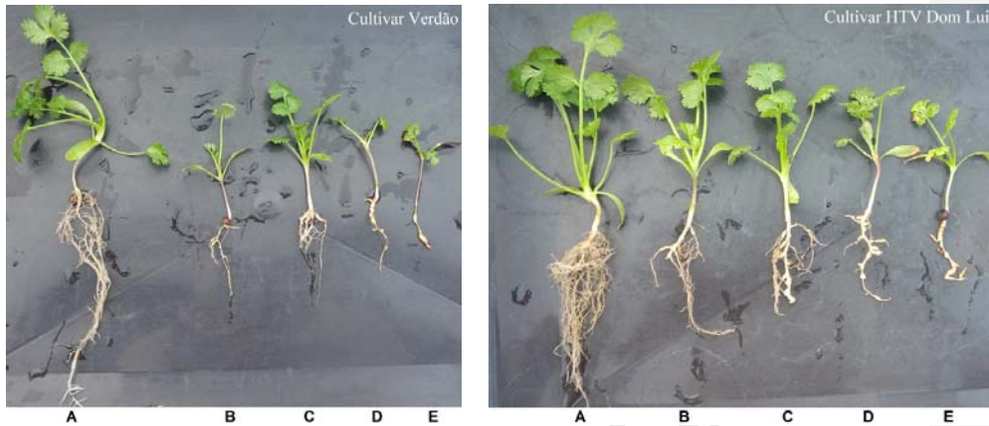
3. RESULTS AND DISCUSSION

157

158 In the treatments containing 16,000 eggs/cell, the plants died within a few days after
 159 germination. As for the treatment with 8,000 eggs/cell, the plants survived until the
 160 transplantation, but they were not able to complete the life cycle, due to the
 161 underdevelopment of both the root system and the shoot (Figure 1).

Comment [EMF2]: ANOVA

162



163

164 **Figure 1.** Coriander plants of Verdão and HTV Dom Luiz, 30 days after inoculation with
 165 different inoculum concentrations of *M. incognita* race 1. A) 0 eggs/plant; B) 1,000
 166 eggs/plant; C) 2,000 eggs/plant; D) 4,000 eggs/plant; E) 8,000 eggs/plant.

167

168 It is possible to observe that with the increase of the concentration of the inoculum there is a
 169 reduction in the development of the root system and the shoot, simultaneously. A similar fact
 170 was observed by [25] in beet cultivars where a linear reduction of vegetative and root
 171 characteristics occurred as the inoculum level of *Meloidogyne incognita*, *M. javanica*, and *M.*
 172 *enterolobii* increased.

173

174 For the variables number of galls in the root system, number of fruits and fruit weight,
 175 interactions between variation sources of variation and concentrations of inoculum X
 176 cultivars were significant at 1% probability. As for the variable number of plants, there were
 177 significant differences only for concentrations of inoculum at 1% of significance. The
 178 coefficients of experimental variation ranged from 12.98% (NGRS) to 27.08% (NF) (Table 2).

179

180 **Table 2.** Summary of the variance analysis of the number of galls in the root system
 181 (NGRS), number of plants (NP), number of fruits (NF) and weight of 100 fruits (WF) in two
 182 cultivars of coriander inoculated with six concentrations of inoculum of *Meloidogyne*
 183 *incognita* race 1.

SV	DF	MS			
		NGRS ⁺	NP	NF ⁺	WF ⁺
Blocks	3	0.09	0.02	2.02	0.002
Concentrations	5	16.29**	2.02**	222.58**	2.79**
Cultivars	1	4.51**	0.02 ^{ns}	330.44**	0.001 ^{ns}
Concentrations*Cultivars	5	0.70**	0.02 ^{ns}	45.96**	0.05**
Error	33	0.07	0.02	3.02	0.01
CV%		12.98	22.35	27.08	13.72

Mean	2.09	0.65	6.41	0.75
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184 † Data transformed by \sqrt{x}
 185 ** Significant at 1% probability
 186 ^{ns} Not significant
 187

188 For the variable NGRS, there was no significant difference between the cultivars only in the
 189 concentrations of 1,000 eggs/plant and 16,000 eggs/plant. In all other concentrations, the
 190 HTV Dom Luiz cultivar presented the highest number of galls compared to the cultivar
 191 Verdão (Table 3).
 192

193 **Table 3.** Scott-Knott grouping test of the number of galls in the root system (NGRS), number
 194 of plants (NP), number of fruits (NF) and weight of 100 fruits (WF) in two coriander cultivar
 195 inoculated with six concentrations of *Meloidogyne incognita* race 1 inoculum, cultivated in
 196 greenhouse.
 197

Variables	Concentration	Cultivars	
		Verdão	HTV Dom Luiz
NGRS	0	-	-
	1,000	10.50 a	10.75 a
	2,000	10.75 a	14.00 b
	4,000	7.00 a	10.75 b
	8,000	2.50 a	6.75 b
	16,000	0.00 a	0.00 a
NP	0	1.00 a	1.00 a
	1,000	1.00 a	1.00 a
	2,000	1.00 a	1.00 a
	4,000	0.75 a	1.00 b
	8,000	0.00 a	0.00 a
	16,000	0.00 a	0.00 a
NF	0	187.25 b	68.25 a
	1,000	259.75 b	67.25 a
	2,000	129.25 b	31.25 a
	4,000	199.00 b	63.75 a
	8,000	0.00 a	0.00 a
	16,000	0.00 a	0.00 a
WF (grams)	0	1.37 a	1.13 a
	1,000	1.36 a	1.11 a
	2,000	1.05 a	1.05 a

4,000	1.52 b	1.01 a
8,000	0.00 a	0.00 a
16,000	0.00 a	0.00 a

198 Means followed by the same lowercase letter in the row did not differ statistically by the Scott-Knott test
 199 at 5% probability.

200

201 In the number of plants that completed the life cycle, in both cultivars, the concentrations of
 202 8,000 and 16,000 eggs/plant did not allow plant survival, thus obtaining the lowest averages
 203 for this variable. For the remaining concentrations, there was no variation within or between
 204 cultivars, except in 4,000 eggs/plant where the cultivar Verdão obtained a lower average
 205 than the cultivar HTV Dom Luiz. In addition, the average was lower than those obtained for
 206 the concentrations of 0, 1,000, and 2,000 eggs/plant. Regarding the number of fruits, in the
 207 concentrations of 0 - 4,000 eggs/plant, Verdão cultivar had a higher number of fruits in
 208 relation to HTV Dom Luiz (Table 3).

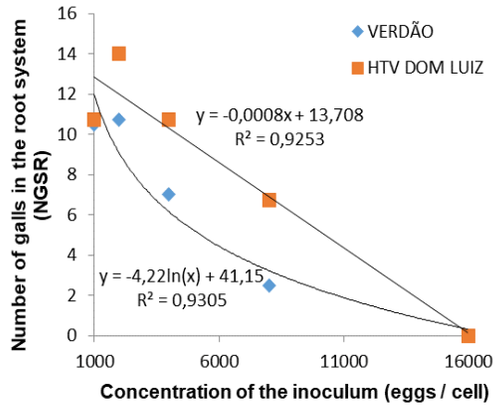
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210 At the inoculum concentrations of 1,000 and 2,000 eggs/plant, both cultivars obtained the
 211 largest number of plants that completed the life cycle, allowing the evaluation of the number
 212 of galls in the root system with subsequent transplantation of the individuals selected for
 213 recombination and attainment of the new improved population.

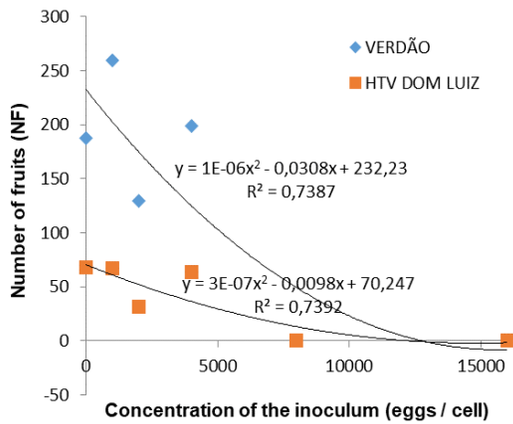
214

215 Based on the obtained regressions, for the variable NGRS, it is noticed that with the
 216 increase of the inoculum concentration there is a reduction of the number of galls in both
 217 cultivars, fact justified by the underdevelopment of the plants under high concentrations of
 218 the nematode. As for the number of fruits, the cultivar Verdão was higher in the
 219 concentration of 1,000 eggs/plant, while with Dom Luiz HTV, the highest yields of fruits were
 220 obtained in concentrations of 0, 1,000 and 4,000 eggs/plant. In both cultivars, there was no
 221 fruit yield in the concentrations of 8,000 and 16,000 eggs/plant, for the plants did not survive
 222 until the reproductive phase. By means of the obtained results, it is observed that the
 223 concentration of 1,000 eggs/plant allows a greater number of fruits for both cultivars (Figure
 224 2).

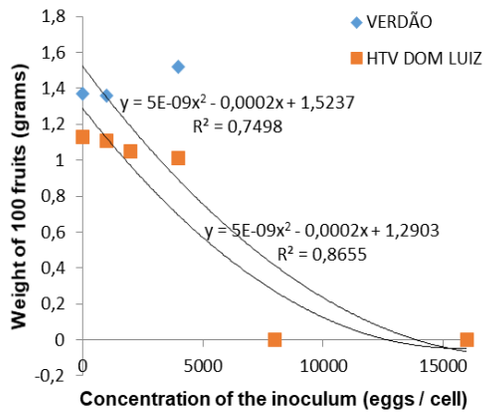
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226



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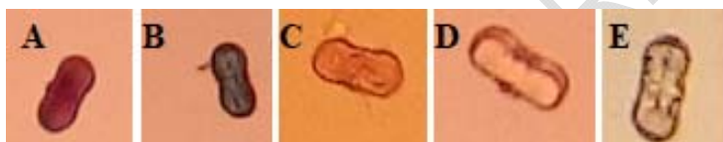
228

229 **Figure 2.** Number of galls in the root system (NGRS), number of fruits (NF) and fruit weight
 230 of Verdão and HTV Dom Luiz cultivars, as a result of different inoculum concentrations.

231 As for the weight of 100 fruits, the highest means obtained in cultivar Verdão were at
232 concentrations 0, 1,000, and 4,000 eggs/plant. In both cultivars Verdão and HTV Dom Luiz,
233 the lowest fruit weight averages were obtained at concentrations of 8,000 and 16,000
234 eggs/plant, due to the death of the plants caused by the intense attack of the pathogen
235 (Figure 2).
236

237 In several crops, the production reduction is reported due to the presence of *M. incognita*, as
238 in the case of soybeans, where losses of 20% to 30% of production are estimated [26].
239 However, in the present work, it was observed that for both the number of fruits and the
240 weight of the 100 fruits, the presence of the pathogen up to the concentration of 4,000
241 eggs/plant did not influence the production and productivity when compared to the control,
242 concentration 0. Possibly, after transplantation to the 2 L pots, plants were able to
243 reestablish the root system and shoot development, not interfering with the flowering and
244 fruit filling. By means of these results, depending on the intensity of the pathogen in a given
245 area of cultivation, it is possible to produce coriander fruits in soils contaminated by *M.*
246 *incognita* race 1.
247

248 As for pollen viability, acetic Carmine and Alexander dyes reacted to the pollen grains, and
249 consequently staining them. As for the tetrazolium solutions, regardless of concentration
250 (1%; 0.75%; 0.50% and 0.25%), they did not stain the pollen at any intensity of red (Figure
251 3).
252



253

254 **Figure 3.** Coriander pollen grains stained with different dyes. Alexander solution, A) viable
255 pollen grain, B) non-viable pollen grain. Acetic Carmine, C) viable pollen grain, D) non-viable
256 pollen grain. Tetrazolium solution, E) non-viable pollen grain.
257

258 The results obtained for the tetrazolium solutions do not corroborate other studies with the
259 same concentrations used in the present study for eggplant [18] and even in concentrations
260 well below as 0.075% in wild passion fruit [27], where tetrazolium was efficient to stain the
261 viable pollens. It is possible that there is some impediment to the penetration of tetrazolium
262 in the structure of the coriander pollen grain, since according to [28], the tetrazolium test is
263 reliable in distinguishing viable and non-viable pollen due to the reaction with the enzyme
264 dehydrogenase of the malic acid that reduces the tetrazolium salt in living tissues where
265 there are ions of H^+ forming a red compound, being related to the cellular respiration of the
266 pollen.
267

268 As fruit yield was verified in the present study, and the plants were managed in a
269 greenhouse where there was no entry of pollinators, it is possible to state that the fruits were
270 obtained from pollination (possibly self-pollination) with the pollens from evaluated plants.
271 Thus, the tetrazolium solutions used were not efficient in distinguishing viable and non-viable
272 pollens in coriander.
273

274 The analysis of variance showed triple interaction (cultivars X inoculum concentrations X
275 dyes) significant at 1% probability. The experimental coefficient of variation was 5.28%
276 (Table 4). The value obtained for the CV% approached that obtained by Santos et al. (2016),
277 which was 3.82%, being considered a low CV% [29].
278

279 **Table 4.** Summary of variance analysis of the percentage of stained pollen grains from two
 280 coriander cultivars inoculated with four inoculum concentrations of *Meloidogyne incognita*
 281 race 1.
 282

SV	DF	MS	
		% of stained pollen grains	
Block	3	26.96	
Genotypes	1	237.62**	
Concentrations	3	95.52*	
Dyes	1	22.56 ^{ns}	
Genotypes*Concentrations	3	171.36**	
Genotypes*Dyes	1	19.51 ^{ns}	
Concentrations*Dyes	3	97.53*	
Genotypes*Concentrations*Dyes	3	181.42**	
Error	45	25.10	
CV (%)		5.28	
Mean		94.82	

283 *Significant at 5% probability

284 **Significant at 1% probability

285 ^{ns} Not significant.

286

287 There were significant differences between the cultivars when developed within
 288 concentrations of inoculum and dyes, only for the acetic Carmine dye at the concentration of
 289 1,000 eggs plant, where the HTV Dom Luiz cultivar had a higher percentage of stained
 290 pollen grains and were, therefore, considered viable (Table 5).
 291

292 **Table 5.** Scott-Knott's grouping test of the percentage of pollen grains stained with two dyes
 293 of two coriander cultivars inoculated with four inoculum concentrations of *Meloidogyne*
 294 *incognita* race 1 grown under greenhouse conditions.
 295

296

Cultivars	Inoculum Concentrations	Dyes	
		Acetic Carmine	Alexander
Verdão	0	96.50 a A β	93.67 a A α
	1,000	75.00 a A α	94.50 b A α
	2,000	97.67 a A β	95.50 a A α
	4,000	97.83 a A β	92.50 a A α
HTV Dom Luiz	0	97.00 a A α	97.00 a A α
	1,000	99.17 a B α	96.33 a A α
	2,000	92.50 a A α	95.67 a A α

4,000

98.16 a A α

98.17 a A α

297 Means followed by the same lowercase letter in the row, upper case for cultivars and Greek for
298 inoculum cultivars in the column do not differ statistically by the Scott-Knott test at 5% probability.
299

300 In the decomposition of concentrations within the cultivars and dyes, only the concentration
301 of 1,000 eggs/plant presented a lower percentage of pollen grains stained in relation to the
302 other concentrations in the cultivar Verdão with acetic Carmine dye.
303

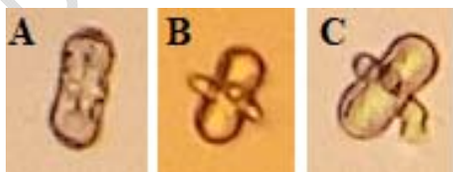
304 As for the dyes, there was variation within the cultivar Verdão at the concentration of 1,000
305 eggs/plant, where acetic Carmine showed a smaller number of grains stained in relation to
306 Alexander. In the study with cane-do-brejo, the lowest values of pollen stained for acetic
307 Carmine and Acid Fuchsin were also observed [7].
308

309 Due to the obtained results, both tested dyes can be used, independently of the
310 concentration of the inoculum used, because the lowest percentage of viable pollen was
311 75% obtained for the cultivar Verdão at the concentration of 1,000 eggs/plant. [30], when
312 working with passion fruit, an allogamous species, consider pollen viability high when values
313 above 70% are obtained, where there is no compromise of breeding work.
314

315 In the evaluation of the pollen in vitro viability, none of the used media allowed pollen
316 germination. The media were tested in the order in which they are found in Table 1. Initially,
317 the means proposed by [31] and [18] were tested. As none of the means presented
318 development of the pollinic tube, mean adaptation was tested in order to verify one that
319 promoted pollen germination, for several factors influence the germination of pollen in vitro
320 from the temperature and period of incubation to the micro and macronutrients in the culture
321 medium, according to [32].
322

323 In the first alteration (C medium), the beginning of the pollen tube elongation was observed
324 after 24 hours of incubation at 25°C. In order to verify if a longer incubation period was
325 necessary, the evaluation was extended for a further 48 hours, observing every 24 hours,
326 totaling 72 hours of observation. However, there was no progression of the pollen
327 elongation, and the length obtained was insufficient to consider the pollen as germinated,
328 considering the criterion proposed by [33] in which the germinated pollen must have a pollen
329 tube length equal to or greater than the diameter of the pollen itself.
330

331 In order to identify an efficient medium in the germination of coriander pollen grains in vitro,
332 adaptations were made in the C medium, varying the concentrations of salts, sucrose, boric
333 acid and phytigel, since according to [34] the types and concentrations of sugar and boron
334 are important in the composition of the culture medium. Although G, I and J media showed
335 the beginning of development of the pollen tube, there was no further growth (Figure 4).
336



342
343 **Figure 4.** Evaluation of the development of the pollen tube cultivated in vitro. A) Absence of
344 tube emission, result obtained for most of the media tested; B) beginning of development of

345 the pollen tube, result obtained in G, I and J media; C) elongation of the pollen tube,
346 observed in C medium.

347

348 Based on the results obtained, there is a need for new studies that seek to identify the factor
349 that is making it impossible to develop the in vitro pollen tube of coriander pollen grains. C
350 medium, presented in this study, can be used as base medium, altering both the sugar and
351 boron as well as the other salt concentrations, and the pH values of the medium. In addition
352 to the concentrations of sugar and boron, pH is a factor that influences the germination of
353 pollen, a fact verified by [31] evaluating media with 3.5 - 6.5 pH in citrus, who verified a linear
354 growth of the number of pollen grains germinated as the pH increased up to the level of 6.5
355 for the cultivar Pêra and Natal. However, in the cultivar Valencia, the best results were
356 obtained at pH around 5.

357

358 5. CONCLUSION

359

360 There is a possibility of inoculation in the egg sowing of 1,000-4,000 eggs/cells of *M.*
361 *incognita* race 1 to evaluate the reaction of coriander genotypes, allowing the quantification
362 of galls and transplantation of the selected plants into pots, directing them to the
363 recombination in open field, since the evaluation based on the number of galls is a non-
364 destructive method and the presence of the pathogen did not compromise the pollen viability
365 and coriander fruit yield in the surviving plants evaluated.

366

367 Seed inoculation and evaluation at 30 days post inoculation provides a 50% reduction of the
368 time required in the selection procedures in which inoculation is performed at 15 days after
369 sowing and evaluation at 45 days post inoculation. In addition, the evaluation through the
370 number of galls - in the early cycles - allows the recombination in the same cycle of the
371 selection, making possible the realization of three selective cycles in a year, thus reducing
372 cost and time to obtain superior genotypes resistant to the nematode.

373

374 Among the dyes used to verify pollen viability, acetic Carmine and Alexander solution were
375 efficient in differentiating viable pollens from nonviable ones, and could be used to verify
376 pollen viability in coriander. None of the tetrazolium concentrations stained the coriander
377 pollen.

378

379 The culture media used for in vitro germination of the pollen grains did not allow the
380 development of the pollen tube, and new studies must be continued in order to adjust a
381 suitable medium for the coriander culture. C medium becomes an option for continuation.

382

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UNDER PEER REVIEW

