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Premature germination of resting spores as a means of protecting brassica crops from *Plasmodiphora brassicae* Wor., (Clubroot)

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7 Running title: Triggering premature spore germination in *Plasmodiophora brassicae*

8 Abstract

9 Clubroot disease causes substantial yield and quality losses in broadacre oil seed and intensive vegetable brassica crops worldwide. The causal microbe *Plasmodiophora brassicae* 10 11 Wor., perennates as soil-borne dormant resting spores. Their germination is triggered by exudates from host roots. A valuable addition to sustainable integrated control strategies 12 could be developed by identifying and synthesising the molecules responsible for stimulating 13 14 resting spore germination. This paper reports experiments in which stimulatory exudates were collected from brassica roots following exposure to infective stages of *P. brassicae*. Analyses 15 identified a germination signalling molecule of circa 1 kDa formed of glucose sub-units. 16 17 Mass spectral analyses showed this to be a complex hexasaccharide carbohydrate with structural similarities to the components of plant cell walls. This is the first report of a host 18 generated hexasaccharides which is capable of stimulating the germination of resting spores 19 of P. brassicae. The implications for environmentally benign control of clubroot are 20 discussed briefly. 21

Keywords: *Plasmodiphora brassicae*, clubroot, resting spores, germination, hexasaccharide,
integrated control

24 Clubroot disease (Plasmodiophora brassicae) causes economically very serious damage to brassica crops worldwide (Dixon, 2009, Strelkov & Dixon, 2014). The disease cycle begins 25 with the germination of environmentally resistant, soil borne resting spores (Dixon 2014). 26 27 This releases biflagellate naked primary zoospores which swim in soil moisture films towards host root hairs (Aist & Williams, 1971). Once inside a host the pathogen reproduces causing 28 disruption of the host metabolism and the development of swollen root tissues. Severely 29 malformed roots lose their normal functions resulting in premature host death. Eventually the 30 roots decay releasing further generations of resting spores into the soil (Dixon. 2006). This 31 32 pathogen is most vulnerable to control strategies during the period from resting spore 33 germination to penetration into host root hairs.

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35 The development of crop protection molecules whose mode of action operates by encouraging resting spore germination offers an effective and sustainable means of control. 36 Root exudates were identified as capable of stimulating resting spore germination by 37 38 Macfarlane, (1970). Subsequently, Craig (1989) showed that root exudates from green broccoli (B. oleracea var. italica) stimulated resting spore germination. Further research 39 demonstrated that the highest levels of resting spore germination (75%) followed treatment 40 with root exudates derived from susceptible cabbages (B. oleracea var. capitata) (Ohi et al, 41 42 2003; Hata et al, 2002). The research reported here describes the isolation and identification 43 of the chemical nature of specific germination stimulators derived from root exudates.

44 Materials and Methods

Heavily galled cabbage (*B. oleracea* var. *capitata*) roots were preserved at -20 °C until required (Dixon, 1976). Resting spore extraction involved defrosting the roots, washing-off residual soil and homogenising portions in 100 ml aliquots of distilled water. The resultant slurries were filtered through four layers of surgical gauze and 44ml of filtrate containing

49 resting spores was decanted as 1.5ml aliquots into Eppendorf tubes. These were centrifuged 50 at 2000 g for 20 minutes. The resultant spore pellets were clarified by repeated suspension in 51 aliquots of 1ml of distilled water and re-centrifuged. Thereafter, the spore pellets were re-52 suspended were in distilled water and stored at -20 °C.

The brassica host used in this research was the clubroot susceptible cabbage (Brassica 53 oleracea var. capitata) cv. Bartolo seed was obtained from a commercial source. When 54 required seed was germinated in Petri plates lined with surgical gauze each moistened with 55 5ml distilled water and held in darkness at 20 °C for two days and then exposed to light. A 56 sample of seedlings was used to determine that the spores of *P. brassicae* obtained by 57 centrifugation and clarification were capable of germination. These were transferred to fresh 58 Petri plates and the roots were sprayed with an aliquot of P. brassicae resting spores at a 59 concentration of 10^7 spores /ml as determine by haemocytometry. After 24 h root samples of 60 61 1 cm length were dissected and placed on a microscope slide mounted in fresh distilled water. This was viewed by microscopy and showed that the resting spores had germinated and 62 63 released primary biflagellate zoospores which were actively swimming around the root samples. This demonstrated that exudates from the roots of cv Bartolo were capable of 64 stimulating the germination of resting spores of *P. brassicae*. 65

Establishing the chemical nature of the triggers of *P. brassicae* resting spore germination coming from cv Bartolo roots required increased volumes of exudates. Seed was sown into a series of ten Petri plates which were prepared as described above. The resultant germinated plants were allowed to grow in the plates placed in light on a north facing laboratory window ledge for 10 days and water in the plates was replenished as required. At the end of this time the seedlings were carefully removed from the surgical gauze and the water squeezed out into a beaker using forceps. Residual moisture was present on the gauze was expressed by placing

73 it in centrifuge tubes and spinning at 2000g for 10 minutes. In total this yielded 42 ml of fluid which contained root exudates produced by the germinating and growing cv Bartolo 74 seedlings. The fluid had a slightly milky appearance it was stored at 4 °C in a laboratory 75 76 refrigerator. It was necessary to demonstrate that this fluid contained root exudates capable of stimulating resting spore germination. Approximately 0.5 ml of fluid was pipetted onto each 77 of five microscope slides and an aliquot of the resting spore suspension was added. Each 78 slide was sealed with nail varnish thereby preventing desiccation. After 20 h examination by 79 microscopy identified motile biflagellate primary zoospores of *P. brassicae* actively 80 81 swimming on the slide. This confirmed the presence of an active compound capable of triggering resting spore germination. 82

Initial experiments with reverse phase high pressure liquid chromatography (HPLC) and 83 elution with several gradients did not find eluates with germination stimulating properties. 84 85 Subsequent experiments with a gravity fed gel filtration column (Sephadex 100; 120 cm x 1cm eluted with 0.1M phosphate buffer) recovered a compound which did stimulate resting 86 87 spore germination. When this stimulant was injected into the column several peaks were obtained in the mass detector trace. The separation was repeated several times using a 88 fraction collector. Fractions of root exudates were collected every 4 minutes. Ultimately 52 89 tubes were collected each filled with 1 ml eluate. The potency of these fractions for 90 stimulating the germination of resting spores of *P. brassicae* was tested. An aliquot of 0.25ml 91 was taken from each fraction and placed on a microscope slide with a similar volume of P. 92 brassicae resting spores suspended in distilled water. The slide was sealed with nail varnish 93 94 and held at room temperature for 20hr. Microscopic examination identified the presence of swimming biflagellate zoospores of P. brassicae. The most active fraction, number 24, 95 stimulated germination in this assay down to 10-fold dilution of the eluates from the fraction 96

97 collector. This positive fraction had a retention time of 85-89 minutes and a molecular weight
98 of 1 KDa. Bradford's reagent tests indicated the absence of proteins. But the large molecular
99 weight suggested that it was a carbohydrate. The active fraction was frozen and freeze-dried
100 to a white residue for mass spectral assays.

Subsequently, more accurate estimates of molecular weight were obtained by HPLC using a 101 Dionex CarboPac MA-1 analytical column (4 x 250mm) with the guard column Dionex 102 CarboPac MA-1 (4 x 50mm) and Mass Detector Sedex model 55. The set-up details were:-103 isocratic gradient A:B (%) 15:85; A contained de-ionised water and B contained 600mM 104 sodium hydroxide, flow rate: 0.4ml/min, temperature was ambient, injection: 20µL, run 105 time was 45 minutes and detection used a Dionex ED40 Electrochemical Detector and 106 mass detector Sedex model 55. The system was calibrated with dextrans of differing 107 molecular weights viz: 2,000,000, 298,000, 9,100 and 8,800 Da in 1 mgml⁻¹ solutions. 108

Samples from the gel column were hydrolysed with 2M hydrochloric acid 1:1 v/v at 100° C 109 110 for 1 hour. Calibration of the column used standard sugar samples, their retention times 111 were: mannose, 20.53; glucose, 22.65 and galactose, 24.97 respectively. The trace from a 1 hr hydrolysis produced a peak in the glucose position. When hydrolysis was extended to 3 112 hrs the peak reached a maximum. Three repetitions of this analysis each produced a single 113 peak in the glucose position. Mass spectral analysis was made on a Liquid 114 Chromatography Quadrupole (LCQ) Classic machine (Thermoscientific, Hemel 115 Hempstead, Hertfordshire) using an electrospray ionization needle voltage of 4.5kV and 116 capillary temperature of 250 °C. The sample was introduced in 0.1% v/v aqueous formic 117 acid and the instrument was scanned between 100 and 2,000 atomic mass units (amu) with 118 a retention time of 2 minutes. The results of mass spectrometry analysis (Figure 1) 119 confirmed that the active compound is a glucose hexamer. A glucose hexamer 120

121 (hexasaccharide) with a molecular weight of 996 Da would have a molecular weight 122 corresponding with that of the compound which stimulated the germination of resting 123 spores of *P. brassicae*. The estimated concentration of the active stimulant of resting spore 124 germination based on the height of the glucose peak in the analysis of hydrolysate was 125 approximately 4μ M.

The LCQ mass spectrum obtained from the active fraction contained several mass peaks in the range from 145 to 1000m/z. The peaks and the range indicated a carbohydrate of 6 hexose units. Several peaks could be interpreted as characteristic of a linear polysaccharide with six glucose units, but the spectrum was more complex than would be anticipated from known fragmentation patterns (Cancilla et al., 1998). It was not possible to deduce the linkage or branching pattern from the data as it is not known if the glucose units are derivatised, with for example N-acetyl groups.

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Results of these experiments confirmed that an active compound capable of stimulating the 134 germination of resting spores of *P. brassicae* and resulting in the release of motile primary 135 zoospores is produced by roots of germinating brassica seedlings. This signalling molecule 136 has now been identified as having a molecular weight of approximately 1 kDa and being 137 composed of glucose subunits forming a hexasaccharide. The hexasaccharide carbohydrate 138 identified in this research has similarities with cell wall polysaccharides (Kiely et al., 2006) 139 which have signalling properties. But literature searches have not found reports associating 140 hexasaccharides with root extracts from hosts of P. brassicae. Nor apparently, is there 141 information regarding the importance of the molecular structure of hexasaccharides and the 142 activation of microbial spore germination. 143

Understanding the chemical nature of signalling molecules in root exudates which elicit 144 resting spore germination permits the development of additional tools for the integrated 145 sustainable control of this pernicious pathogen. Currently there are no agrochemicals 146 available for the control of *P. brassicae*, resistance genes are of limited occurrence and their 147 usefulness may be eroded quite quickly by the appearance of tolerant physiological races 148 (Dixon, 2014). Control strategies rely on soil husbandry, crop nutritional and rotational 149 techniques. Individually none of these methods offers complete control, mitigation of damage 150 sufficient for the culture of profitable crops comes solely from combinations of the 151 152 techniques. Adding further elements into integrated control strategies increases their effectiveness and longevity as described by Rashid et al (2013). The research reported here 153 offers a further dimension for control through the development of hexassacharide molecules 154 155 into formulations which might be applied as soil or transplant applied treatments which diminish the inoculum potential of *P. brassicae* populations in infested soil. Also there are 156 opportunities for use hexasaccharide formulations in the absence of susceptible crops thereby 157 cleaning up land in advance of growing brassica crops. 158

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213	Figure 1

- Mass Spectrum of the hexasaccharide which stimulated germination of resting spores of *P. brassicae*

