1	Low doses of selenium nanoparticles enhance the performance of the				
2	generalist parasitoid Anisopteromalus calandrae				
3					
4	RUNNING TITLE: SELENIUM NANOPARTICLES AND A PARASITOID				
5					
6	James Rudolph Miksanek ¹ *, Charles Adarkwah ^{1,2,3} , Midori Tuda ¹ *				
7					
8	¹ Laboratory of Insect Natural Enemies, Institute of Biological Control, Faculty of Agriculture, Kyushu				
9	University, Fukuoka 819-0395, Japan				
10					
11	² Department of Horticulture and Crop Production, School of Agriculture and Technology, Dormaa-				
12	Ahenkro Campus, University of Energy and Natural Resources, Sunyani, Ghana				
13					
14	³ Humboldt-University of Berlin, Division Urban Plant Ecophysiology, Faculty Life Sciences, 14195,				
15	Berlin, Germany				
16					
17	*Correspondence:				
18	Laboratory of Insect Natural Enemies				
19	Faculty of Agriculture				
20	Kyushu University				
21	Fukuoka 819-0395 (Japan)				
22	email: miks0007@umn.edu				

23 ABSTRACT

24 BACKGROUND: The environmental and economic costs of conventional insecticides have stirred an

25 interest in alternative management tactics, including the use of nanotechnologies. Selenium nanoparticles

26 (SeNPs) have many applications in agriculture but may not be compatible with biological control;

27 however, low concentrations of SeNPs may benefit natural enemies via hormesis. This study investigates

28 the dose-dependent effects of SeNPs $(0-1000 \text{ mg L}^{-1})$ on Anisopteromalus calandrae (Howard)

29 (Hymenoptera: Pteromalidae), a generalist parasitoid of stored product pests.

30

31 RESULTS: The LC₅₀ of SeNPs was 1540 mg L^{-1} for female parasitoids and 1164 mg L^{-1} for males

32 (which was not significantly different between sexes). SeNPs had a significant hormetic effect; average

lifespan increased by 10% at a dose of 4.03 mg L^{-1} for females and by 35% at 13.83 mg L^{-1} for males. In

34 a bioassay including hosts, low-dose SeNPs (25 mg L^{-1}) enhanced female performance; lifespan

35 increased by 23% and the number of offspring increased by 88%. However, the number of emerging hosts

36 (azuki bean beetle *Callosobruchus chinensis* (L.) (Coleoptera: Chrysomelidae: Bruchinae)) did not

37 significantly decrease; in the absence of parasitism, SeNPs improved emergence by 17%.

38

39 CONCLUSION: Because higher concentrations of SeNPs reduced parasitoid lifespan, whereas low doses

40 enhanced not only parasitoid performance but also host emergence, practitioners will need to pay special

41 attention when optimizing SeNP formulations to maximize their contribution to pest control or in the

42 mass rearing of insect parasitoids.

43

Keywords: integrated pest management, biological control, nanotechnology, biological control, stored
 product pests, dose-response, hormesis

46 1. INTRODUCTION

47 The complementarity of chemical, cultural, biological, and other control strategies is a key aspect of

- 48 integrated pest management (IPM)^{1,2}. Because there tend to be compatibility issues between conventional
- 49 insecticides and natural enemies $^{3-5}$, and due to the environmental and economic costs of these
- 50 pesticides^{6,7}, there are ongoing efforts to counter an overreliance on chemical control and incorporate
- 51 safer biorational alternatives to improve management outcomes and meet sustainable development goals
- 52 $(SDGs)^{8-10}$. Emerging research is also investigating the possible roles of nanotechnology in pest
- 53 management, including the use of entomotoxic nanoparticles (NPs) or nano-encapsulated pesticides^{11,12};
- 54 despite this effort, the ecological risks of NPs remain relatively unknown^{11,13–15}. Due to the nonspecific
- 55 mode of action of elemental (or elemental oxide) NPs¹⁶, these products might pose a risk towards
- 56 nontarget organisms, including natural enemies.

57 Even though the sublethal effects of insecticides on natural enemies are typically negative^{3,17}, there are also instances in which they can be positive¹⁸. The term "hormesis" (or "hormoligosis" in 58 59 insects¹⁹) refers to the positive or stimulatory effect of low doses of a chemical stressor, which can be characterized by a biphasic dose-response curve²⁰. Since pesticide residues are often present throughout 60 the agroecological landscape, it is important to understand how these (and other stressors) influence the 61 62 biological control services provided by natural enemies. A hormetic response may improve pest 63 suppression or be applicable to the mass rearing and release of commercial biological control agents^{18,21}. 64 Although there are many variables that affect the persistence of anthropogenic NPs in the environment (and their relationship to naturally occurring NPs)^{15,21,22}, relatively small or residual amounts of NPs 65

66 might offer a direct benefit to biological control agents by hormesis.

67 *Anisopteromalus calandrae* (Howard) (Hymenoptera: Pteromalidae) is a cosmopolitan parasitoid 68 that attacks a broad range of hosts, including the azuki bean beetle *Callosobruchus chinensis* (L.) and the 69 cowpea seed beetle *C. maculatus* (F.) (Coleoptera: Chrysomelidae: Bruchinae)²³. As solitary idiobiont

- 70 ectoparasitoids, females typically lay one egg after paralyzing a host²⁴ (usually a late-instar larva or pupa),
- and the developing parasitoid larva feeds from outside of its host; additional parasitoid-induced mortality

can arise via host feeding²⁵, a process through which adult female *A. calandrae* attack and consume the A_{12}

hemolymph of hosts (preferably younger instars²⁵) to acquire nutrients for egg maturation²⁵⁻²⁷ (and, when

- feeding on *C. chinensis*, to enhance longevity as well; JRM, personal observation). Because of the
- importance of A. calandrae in the control of bruchine beetles (and other coleopteran pests)^{28–32}, it is
- 76 important to test their compatibility with the entomotoxic NPs that have garnered recent attention—
- selenium nanoparticles (SeNPs), in particular^{33,34}—as a possible new control strategy for these insects.
- SeNPs have several potential applications in agriculture: promoting plant growth, bolstering plant
 defenses, and acting as an antimicrobial, nematocidal, or insecticidal agent, depending on the

- concentration or formulation $^{35-37}$. The entomotoxic effects of SeNPs arise through the slow release of 80 81 elemental selenium, which, depending on the route of entry, may amass in the Malpighian tubules and midgut, negatively impacting the growth, development, and survival of the target insect³⁸⁻⁴⁰. While SeNPs 82 have shown some promise as an alternative control method for bruchine beetles, including C. 83 84 chinensis^{33,34}, there is evidence they may also harm the parasitoid A. calandrae when applied 85 concurrently³⁴. However, lower doses of SeNPs may actually enhance parasitoid performance via 86 hormesis. The purpose of the present study is to investigate the dose-dependent effects of SeNPs (0-1000 87 mg L^{-1}) on the lifespan of the generalist parasitoid A. calandrae and to test the hormetic effects of low 88 doses of SeNPs (25 mg L^{-1}) on parasitoid performance (lifespan, fecundity, offspring emergence time, 89 age-dependent parasitism rate, and host suppression), all of which is relevant in the control of the azuki 90 bean beetle and has important implications for other IPM programs that incorporate A. calandrae or 91 related generalist parasitoids.
- 92

93 **2. METHODS**

- 94 Colonies of the azuki bean beetle, *C. chinensis* (strain jC, originating from Japan), and its parasitoid *A*.
- 95 calandrae (also from Japan) have been maintained under laboratory conditions (30°C, 60% R.H., 16:8
- 96 L:D) for more than 20 years on dried azuki beans, Vigna angularis var. angularis (Willdenow) Ohwi &
- 97 Ohashi (Fabaceae) (Daiwa Grain Co., Obihiro, Japan). The chemical synthesis of SeNPs^{41,42} (amorphous
- 98 and approximately 5–10 and 60–100 nm in diameter) was previously described by Miksanek & Tuda³³; in
- brief, a 20-mL solution of SeNPs (1000 mg L^{-1}) was prepared by reducing sodium selenite (43.8 mg) with
- 100 ascorbic acid (2 mL, 0.633 M) in ultrapure water (17.9 mL; Milli-Q, 18.2 MΩ·cm) and stabilizing with
- 101 polysorbate 20 (100 μ L). Additional SeNP concentrations of 1, 10, 25, 50, 100, 200, and 500 mg L⁻¹ were
- 102 made by diluting the original solution with ultrapure water; a 0 mg L^{-1} control solution (containing only
- 103 ultrapure water) was also prepared.
- 104

105 **2.1 Dose-dependent toxicity**

106 2.1.1 Bioassay

- 107 Newly emerged (< 24 h) adult parasitoids (males and females) were placed individually in mini-sized
- 108 Petri dishes (35 mm in diameter, 10 mm in height) before adding a 20-µL droplet of SeNPs (0, 1, 10, 50,
- 109 100, 200, 500, or 1000 mg L^{-1} (0–20 µg SeNPs)). Each Petri dish was gently agitated to evenly coat all
- 110 surfaces as well as the parasitoid with SeNPs. After this initial treatment, the parasitoids were placed in a
- 111 climate-controlled chamber (30°C, 60% R.H., 16:8 L:D) and monitored daily until death. All eight SeNP
- 112 concentrations were replicated 25 times for each sex, for a total of 400 parasitoids.
- 113

114 2.1.1 Analysis

115 The concentration-dependent effects of SeNPs on parasitoid lifespan (age-at-death, x) were described

- 116 using the Cedergreen-Ritz-Streibig modification of a log-logistic model, which allows for an effect of
- 117 hormesis 43,44 . This model takes the form:
- 118

$$f(x) = \frac{d + f\exp(-1/x^{\alpha})}{1 + \exp(b(\log(x) - \log(e)))}$$
(1)

119

120 in which d is the upper horizontal asymptote and f is the magnitude of the effect of hormesis (with α

121 affecting the shape of the peak); the parameters *b* and *e* have no direct biological interpretation in the

122 Cedergreen-Ritz-Streibig modification of the log-logistic model (in the related three-parameter log-

logistic model, b is a scaling factor and e is an inflection point that corresponds with the EC_{50} or

- 124 LC_{50})^{43,44}. If the parameter *f* is equal to 0, then there is no hormetic effect and the model simplifies to a
- 125 three-parameter log-logistic curve 43,44 .
- 126 Model fitting and analysis was conducted in R version 4.2.2. The term α was set to 0.5 when 127 fitting the four-parameter Cedergreen–Ritz–Streibig model^{43,44}. Parasitoid sex was included as a grouping
- 128 factor when estimating the parameters *b* and *d* (the model was simplified because preliminary analyses
- 129 showed no significant effect of sex on estimates of *e* and *f*). The package *drc* was used in the analysis,
- 130 which also allowed for estimating the maximum effect of hormesis from the mean response predicted by
- 131 the model (using the "MAX" function⁴⁴, as described by Cedergreen et al.⁴³), lethal concentrations (e.g.
- 132 LC₅₀), and calculating *t*-statistics for comparisons between sexes^{43,44}.
- 133

134 **2.2 Low-dose SeNPs and parasitoid performance**

135 2.2.1 Bioassay

136 The effects of a low dose of SeNPs on the lifespan and reproductive success of *A. calandrae*, as well as

- 137 offspring emergence timeline, were tested by exposing parasitoids to infested azuki beans treated with
- 138 either SeNPs or water. The experimental design consisted of a full factorial for parasitism (A. calandrae
- present or absent) and SeNP application (0 or 25 mg L^{-1} SeNPs). For this bioassay, a concentration of 25
- 140 mg L^{-1} SeNPs was chosen because a preliminary analysis of dose-dependent survivorship suggested that
- 141 this approximate concentration may best enhance parasitoid performance.
- 142 Adult parasitoids were collected from a laboratory colony within 24 h of emergence. Next, each
- 143 male-female pair was treated with a 20-µL droplet of SeNPs (0.5 µg SeNPs) or ultrapure water (as
- 144 previously described) and provided 8 azuki beans (1.1 g) treated concurrently and in the same manner (by
- 145 gently agitating each dish evenly coat all surfaces as well as the beans with SeNPs) with a $30-\mu$ L droplet³³

146 (0.75 µg SeNPs). Each bean was infested by eight mixed-stage C. chinensis larvae (indirectly counted as 147 the number of eggs on the surface of the bean); by the time of parasitoid exposure, most larvae were 15– 148 16 days old, which is the favored host stage for parasitism; the smaller number of younger larvae are the 149 preferred size for host-feeding. Beans and parasitoids were placed in a small Petri dish (55 mm in 150 diameter, 15 mm in height). The male parasitoid was discarded after 24 h to prevent excessive sexual 151 harassment. Parasitoid mortality was monitored at 24-h intervals. After four days, females were provided 152 with a fresh set of infested beans (of the same treatment conditions), and the original sets of beans were 153 checked daily to record the number of emerging adults (hosts and parasitoids). The second set of beans 154 were not monitored for daily emergence, but the total number of emerged parasitoids was recorded three 155 weeks after the death of the parental parasitoid in order to calculate lifetime reproduction. Each 156 combination of treatments was replicated 30 times, for a total of 120 replicates; daily emergence was only 157 recorded from a subset of replicates (24 replicates for each of the treatments in the absence of parasitism 158 and 25 from each of those with a parasitoid present).

159

160 2.2.2 Analysis

The effects of SeNPs on the lifespan and fecundity of adult female parasitoids were tested with a Cox 161 162 proportional hazards model and linear regression model, respectively. A binomial GLM was used to test 163 the effects of SeNP treatment on the sex ratio of the resulting parasitoid offspring. The effects of SeNPs 164 on the development time of offspring (days to adult emergence) were assessed with a Cox proportional 165 hazards model, with sex and the interaction between sex and SeNP treatment as additional factors; the 166 identity of the maternal parasitoid was included as a random effect by clustering.

167 To further investigate the reproduction of female parasitoids, a linear regression was used to 168 assess the relationship between the log-transformed early rate of reproduction (the age-corrected number 169 of offspring per day, calculated as the number of offspring produced from days 0–3 (inclusive) divided by 170 the lifespan of the parasitoid) and SeNP treatment on the reproductive rate later in life (from day 4-death, 171 calculated in the same fashion); only parasitoids with a lifespan greater than 4 days and a nonzero early 172 reproductive rate were included in this analysis. Next, to investigate the influence of SeNPs on the 173 lifetime reproductive success (total number of offspring) of female parasitoids as a function of age, an 174 asymptotic Michaelis-Menten-type model was fit to the data. This model predicts the total number of 175 offspring of a female parasitoid living to age *t* as: 176

$$f(t) = \frac{m(t - t_{\rm pre})}{\beta \exp(-\gamma [\rm Se]) + t - t_{\rm pre}}$$
(2)

178 Analogous to the standard Michaelis-Menten model, the parameter *m* denotes the upper asymptote of the

- 179 saturating curve and β is associated with the midpoint of the saturating curve; biologically, *m* represents
- 180 the maximum number of parasitoid offspring (limited by host depletion or as a more general effect of
- 181 parasitoid senescence). The modifications on the Michaelis-Menten model include an x-intercept (t_{pre}) and
- 182 the effect of SeNPs (exp($-\gamma$ [Se])). The term t_{pre} applies a horizontal transformation to the curve;
- 183 biologically, *t*_{pre} represents the length of the pre-oviposition period of the adult female (a female might
- 184 use this time for mating, sperm storage, host feeding, or egg maturation). The function $exp(-\gamma[Se])$
- applies an effect γ of SeNPs ([Se] = 0 or 25 mg L⁻¹) to β ; thus, at a concentration of 0 mg L⁻¹ (or if there
- 186 is no effect γ), then β is unaffected. For simplicity, the Michaelis–Menten-type model only considers the
- 187 effect of SeNPs on the parameter β ; however, this formulation was also supported by an informal analysis
- 188 of the shortest- and longest-lived parasitoids (0–3 and 9–12 days, respectively), in which were there no
- 189 differences in the reproduction of control and SeNP-treated parasitoids in either age class, suggesting that
- 190 m and t_{pre} are likely independent of SeNP treatment.
- Finally, the effects of SeNPs and parasitism on the number and sex ratio of emerging hosts were tested with a linear mixed-effects model (LMM) and binomial generalized mixed-effects model (GLMM), respectively. In both models, the identity of the source colony was included as a random effect to account
- 194 for potential variation in the stage structure or exact number of hosts; because source colony did not
- 195 improve the statistical models of parasitoid life history (described above), this random effect was only
- 196 included in these analyses of the host population.
- 197All modeling and statistical analyses were conducted in R version 4.2.2. The Michaelis–Menten-198type model was fit using nonlinear least squares; confidence intervals for model predictions were199calculated by Monte Carlo simulation (n = 100000). The packages *car*, *lme4*, *merTools*, *survival*, *coxed*,
- 200 and *propagate* were also used in the analyses.
- 201

202 **3. RESULTS**

203 **3.1 Dose-dependent toxicity**

- 204 The lifespan of adult parasitoids was measured in response to a range of SeNP concentrations (0–1000 mg
- 205 L⁻¹). In fitting the Cedergreen–Ritz–Streibig-modified log-logistic model to the concentration-dependent
- effects of SeNPs on the lifespan of male (n = 200) and female (n = 200) parasitoids (with parasitoid sex as
- a grouping factor when estimating b and d), the estimates for all parameters were significant (Figure 1,
- 208 Table 1). The upper asymptote *d*—which represents the maximum lifespan in the absence of SeNPs—was
- 4.55 ± 0.15 d (estimate \pm SE) for females and 3.01 ± 0.16 d for males, and the difference between sexes
- was significant (t = 6.27, p < 0.001) (Figure 1; Table 1); the estimates of parameter b (a scaling factor)
- 211 were marginally different between sexes as well (t = -1.91, p = 0.057) (Table 1).

The LC₅₀ was 1540 ± 391 mg L⁻¹ for females (estimate \pm SE) and 1164 ± 224 mg L⁻¹ for males, but there was no significant difference in the estimates of LC₅₀ between sexes (t = 1.19, p = 0.237). For females, the average age at death of SeNP-treated individuals was 5.02 d at a concentration of 4.03 mg L⁻¹; for males, the average age at death of SeNP-treated individuals was 4.06 d at a concentration of 13.83 mg L⁻¹ (Figure 1; Table 1).

217

218 **3.2 Low-dose SeNPs and parasitoid performance**

219 Parasitoid performance was assessed in response to a low dose (25 mg L^{-1}) of SeNPs to investigate the 220 effects of hormesis on a generalist biological control agent. The performance parameters that were 221 measured included parasitoid lifespan and fecundity, offspring sex ratio and development time, parasitoid

- 222 reproductive rate and lifetime reproductive success, and host emergence and sex ratio.
- Adult female parasitoids (n = 60) treated with 25 mg L⁻¹ SeNPs lived significantly longer than control parasitoids; adult lifespan was 7.53 ± 0.52 and 6.10 ± 0.48 d, respectively (mean \pm SE) (Figure 2a; Table 2). SeNP-treated females also produced a greater number of offspring; the total number of offspring was 13.0 ± 1.1 vs. 6.9 ± 1.1 (Figure 2b; Table 2). The sex ratio (n = 56) and development time (n = 285) of the offspring of SeNP-treated parasitoids did not differ from those of the control (Table 2). Regardless of SeNP treatment, female offspring (n = 233) emerged later than males (n = 52); the time to emergence was 13.66 ± 0.09 and 12.73 ± 0.18 d, respectively (Table 2).
- For ovipositing female parasitoids with an age of death > 4 d (n = 47), the early reproductive rate 230 231 was inversely correlated with the rate of reproduction later in life (offspring per day, days 4-death) ($F_{3,43}$ = 14.94, multiple R^2 = 0.510, p < 0.001) (Figure 3a). As the early rate of reproduction increased, there 232 was a decrease in the rate of reproduction later in life; treatment with 25 mg L^{-1} SeNPs increased the rate 233 234 of reproduction overall (Figure 3a; Table 3a). To investigate the lifetime reproductive success of all 235 female parasitoids (n = 60), a Michaelis–Menten-type model was implemented, in which all parameter 236 estimates were significant (Figure 3b; Table 3b). The maximum number of offspring m was 44.7 ± 17.8 237 (estimate \pm SE) and the preoviposition period was 1.90 ± 0.30 d (Table 3b). There was also a significant effect of SeNP treatment γ , with 25 mg L⁻¹ SeNPs increasing the lifetime reproductive success of female 238 239 parasitoids (Figures 3a and 3b; Tables 3a and 3b).
- Finally, the effects of SeNPs and parasitism on the host population were evaluated as the number and sex ratio of the emerging hosts in each Petri dish (n = 120). The number of emerging hosts was lower in the presence of a parasitoid than in the absence of the parasitoid but, when treated with SeNPs, was higher in the absence of parasitism (Figure 4a, Table 4). When treated with 0 mg L⁻¹ SeNPs, there were an estimated 21.5 (95% prediction interval: 15.0–28.8) emerging hosts per Petri dish in the absence of parasitism and 19.1 (12.3–26.1) emerging hosts per dish in the presence of a female parasitoid; when
 - 8

- treated with 25 mg L^{-1} SeNPs, there were 25.1 (18.6–31.4) emerging hosts per dish in the absence of
- parasitism and 17.1 (10.9–23.4) emerging hosts with a parasitoid present (Figure 4a, Table 4). Finally, the
- sex ratio of the emerging hosts shifted from a male bias (0.54 [0.51-0.57]) at 0 mg L⁻¹ SeNPs to a slight
- female bias (0.47 [0.45–0.51]) when treated with 25 mg L^{-1} SeNPs (Figure 4b, Table 4).
- 250

251 4. DISCUSSION

- 252 Because of the environmental and economic costs of conventional chemical insecticides, there is ongoing
- 253 interest in exploring alternative management tactics, including the use of nanotechnologies. Selenium
- 254 nanoparticles (SeNPs) have many applications in agriculture, but their nonspecific entomotoxic mode of
- action may limit their compatibility with biological control agents; however, low concentrations may
- benefit natural enemies via hormesis. The present study found that, for the generalist parasitoid A.
- 257 *calandrae*, the LC₅₀ of SeNPs was 1.5 g L^{-1} for female parasitoids and 1.2 g L^{-1} for males, and an
- analysis of the dose-response curves identified a significant hormetic effect; for females, the average age
- at death increased by 10% at a dose of 4.03 mg L^{-1} and, for males, increased by 35% at 13.83 mg L^{-1} . In a
- 260 follow-up bioassay that provided female parasitoids with hosts, low-dose SeNPs (25 mg L^{-1}) were found
- to enhance parasitoid performance; female lifespan increased by 23% and the number of offspring
- 262 increased by 88%. Regardless, the number of emerging azuki bean beetles (the host *C. calandrae*) did not
- significantly decrease after treatment with low-dose SeNPs but rather, in the absence of parasitism,
- increased by 17%.

265 There exists a delicate balance in the essential role of selenium as a micronutrient and its dosedependent toxicity in both vertebrate and invertebrate animals^{35–40,45–49}. In insects, diets with supplemental 266 selenium can benefit growth, survival, and reproduction, but higher concentrations can have the opposite 267 effect^{36,38-40,46}. While insects play a role in the biogeochemical cycling of selenium^{47,49}, little is known of 268 269 the impact of selenium on predator-prey or host-parasitoid ecology; the biotransfer or biomagnification⁵⁰ of selenium can negatively affect the development and fitness of predators⁵¹, and higher concentrations of 270 271 selenium can potentially disrupt host-parasitoid population dynamics, with Cotesia marginiventris 272 (Cresson) (Hymenoptera: Braconidae) and the beet armyworm, Spodoptera exigua (Hubner)

- 273 (Lepidoptera: Noctuidae), serving as an example^{49,52}. Even though SeNPs have a lower toxicity and
- higher bioavailability than inorganic or organic forms of selenium³⁷, applications of SeNPs ranging in
- 275 concentration from 100–500 mg L^{-1} can still have significant entomotoxic effects^{37,53}, including on the
- azuki bean beetle^{33,34} (although its endosymbiont *Wolbachia* can mediate resistance³³).
- In the present study of the *C. chinensis–A. calandrae* system, the parasitoid *A. calandrae* also began showing adverse effects within the 100–500 mg L^{-1} range^{33,34}. Thus, the concurrent use of SeNPs and biological control would not be compatible in this case. Similarly, since the pest species also appears

to exhibit a hormetic response to low-dose SeNPs (25 mg L^{-1}), this limits the complementary of SeNPs 280 281 and biological control, at least in the short term; additional studies that extend beyond the single-282 parasitoid, single-generation paradigm (and that also address the timing of SeNP applications and residual 283 effects), would be required to identify any long-term consequences that might be relevant for pest 284 suppression. But first, there is the fundamental question of whether the performance of A. calandrae is 285 enhanced as a result of a response to SeNPs as a chemical stressor (i.e. hormoligosis/hormesis), via the 286 potential benefits of selenium as a micronutrient (parasitoids could be observed drinking from dispersed 287 droplets of SeNPs [or water] before the application completely dried, although this would not offer the 288 same benefits as supplemental honey or $sugar^{26,54}$), or through some combination of the two. The positive effects of low-dose SeNPs (25 mg L^{-1}) on lifespan (and fecundity) were more 289 290 prominent when hosts were available for host feeding and oviposition, which suggests that SeNPs may at 291 least alter the nutritional physiology of female parasitoids (and, if linked to host feeding, this may introduce an additional source of density dependence in the system^{55,56}). However, because selenoproteins 292 293 have largely been lost in the related parasitoid Nasonia vitripennis (Walker) (Hymenoptera: Pteromalidae) 294 and the honey bee Apis mellifera L. (Hymenoptera: Apidae)^{57,58}, the positive effects of low-dose SeNPs 295 are more likely to be a hormetic response to small amounts of oxidative stress or via a reduced stress threshold rather than as a positive response to an essential micronutrient $^{69-61}$. (Note that most Coleoptera 296 also lack selenoproteins^{57,58,61} as well, so a hymenopteran parasitoid and its coleopteran host might be 297 298 expected to exhibit similar physiological responses to SeNPs, which could complicate control efforts.) 299 Regardless, the ultimate result is an enhancement of parasitoid performance—at least for the treated 300 parasitoids, since any effects on subsequent generations are currently unknown. So, while not a

301 complementary control strategy on its own, and offering no net gain as a tactic for enhancing parasitism

302 over a single generation in the field, the use of low-dose SeNPs (in conjunction with supplemental

nutrients, although this would require further study) may instead have more promising applications in the

mass rearing of commercial control agents for inoculative or inundative release in augmentation
 biological control¹⁸.

306

307 **5. CONCLUSION**

308 Overall, this study provides insight into the role of SeNPs in the control of the azuki bean beetle and has 309 important implications for other programs that incorporate the generalist *A. calandrae* or closely related

310 parasitoids; there are also potential applications in the mass rearing of biological control agents. The

- 311 results of the present study indicate that, because higher concentrations of SeNPs reduced parasitoid
- 312 lifespan—whereas a low-dose treatment benefitted not only the parasitoid but the host as well—the
- 313 effects of SeNPs and biological control may neither be compatible nor complementary tactics. However,

- 314 future studies on the trade-offs between lifespan and reproduction as well as on population-level
- 315 responses (extending beyond single-parasitoid, single-generation experiments) may offer additional
- 316 insight into the possible roles of SeNPs in the field of pest control.

317 6. ACKNOWLEDGEMENTS

318 **6.1 Author contributions**

- 319 All authors contributed to the study conception and design. Material preparation and data collection were
- 320 performed by JRM and CA. JRM and MT participated in data analysis. The first draft of the manuscript
- 321 was written by JRM, and all authors commented on subsequent versions of the manuscript. All authors
- 322 read and approved the final manuscript.
- 323

324 6.2 Data availability

- 325 The datasets and R script associated with the current study are available from the primary corresponding
- author on request.
- 327

6.3 Funding

- 329 JRM is an International Research Fellow of Japan Society for the Promotion of Science (JSPS). CA was
- 330 supported by an Invitational Fellowship from JSPS. MT received additional support by KAKENHI
- 331 (19K06840) from JSPS.
- 332

333 7. CONFLICT OF INTEREST STATEMENT

- 334 The authors declare no conflicts of interest and affirm that the present study was carried out in adherence
- to all applicable ethical standards.

336 8. REFERENCES

- Kogan M, Integrated pest management: Historical perspectives and contemporary developments.
 Annu Rev Entomol 43:243–270 (1998).
- Farrar JJ, Ellsworth PC, Sisco R, Baur ME, Crump A, Fournier AJ, et al., Assessing the compatibility
 of a pesticide in an IPM program. *J Integr Pest Manag* 9:3 (2018).
- 341 3. Desneux N, Decourtye A, Delpuech JM, The sublethal effects of pesticides on beneficial arthropods.
 342 Annu Rev Entomol 52:81–106 (2007).
- 343 4. Beers EH, Mills NJ, Shearer PW, Horton DR, Milickzy ER, Amarasekare KG, et al., Nontarget
- effects of orchard pesticides on natural enemies: Lessons from the field and laboratory. *Biol Control* 102:44–52 (2016).
- Schmidt-Jeffris RA, Beers EH, Sater C, Meta-analysis and review of pesticide non-target effects on
 phytoseiids, key biological control agents. *Pest Manag Sci* 77:4848–4862 (2021).
- Bernard B. Pimentel D, Environmental and economic costs of the application of pesticides primarily in the
 United States. *Environ Dev Sustain* 7:229–252 (2005).
- Damalas CA, Eleftherohorinos IG, Pesticide exposure, safety issues, and risk assessment indicators.
 Int J Environ Res Public Health 8:1402–1419 (2011).
- Rosell G, Quero C, Coll J, Guerrero A, Biorational insecticides in pest management. *J Pestic Sci* 33:103–121 (2008).
- Dangles O, Casas J, Ecosystem services provided by insects for achieving sustainable development
 goals. *Ecosyst Serv* 35:109–115 (2019).
- 356 10. Struelens Q, Silvie P, Orienting insecticide research in the tropics to meet the sustainable
- development goals. *Curr Opin Insect Sci* 40:24–30 (2020).
- 11. Kah M, Hofmann T, Nanopesticide research: Current trends and future priorities. *Environ Int* 63:224–
 235 (2014).
- Athanassiou CG, Kavallieratos NG, Benelli G, Losic D, Usha Rani P, Desneux N, Nanoparticles for
 pest control: Current status and future perspectives. *J Pest Sci* 91:1–15 (2018).
- 13. Crane M, Handy RD, Garrod J, Owen R, Ecotoxicity test methods and environmental hazard
 assessment for engineered nanoparticles. *Ecotoxicology* 17:421–437 (2008).
- 14. Hansen SF, Michelson ES, Kamper A, Borling P, Stuer-Lauridsen F, Baun A, Categorization
 framework to aid exposure assessment of nanomaterials in consumer products. *Ecotoxicology* 17:438–447 (2008).
- 367 15. Wagner S, Gondikas A, Neubauer E, Hofmann T, von der Kammer F, Spot the difference:
- 368 Engineered and natural nanoparticles in the environment—release, behavior, and fate. *Angew Chem*
- 369 *Int Ed* 53:12398–12419 (2014).

16. Benelli G, Mode of action of nanoparticles against insects. *Environ Sci Pollut Res* 25:12329–12341

371 (2018).

- 372 17. Stark JD, Vargas R, Banks JE, Incorporating ecologically relevant measures of pesticide effect for
 373 estimating the compatibility of pesticides and biocontrol agents. *J Econ Entomol* 100:1027–1032
 374 (2007).
- 18. Cutler GC, Amichot M, Benelli G, Guedes RNC, Qu Y, Rix RR, et al., Hormesis and insects: Effects
 and interactions in agroecosystems. *Sci Total Environ* 825:153899 (2022).
- 377 19. Luckey TD, Insecticide hormoligosis. *J Econ Entomol* 61:7–12 (1968).
- 20. Cutler GC, Insects, insecticides, and hormesis: Evidence and considerations for study. *Dose– Response* 11:154–177 (2013).
- 380 21. Guedes RNC, Cutler GC, Insecticide-induced hormesis and arthropod pest management. *Pest Manag* 381 *Sci* 70:690–697 (2013).
- 22. Malakar A, Kanel SR, Ray C, Snow DD, Nadagouda MN, Nanomaterials in the environment, human
 exposure pathway, and health effects: A review. *Sci Total Environ* 759:143470 (2021).
- 23. Zhang Y, Yu F, Wu L, Dai R, Yang H, Zhang X, Hu D, Life history traits of the parasitoid
 Anisopteromalus calandrae (Hymenoptera: Pteromalidae) on three beetle hosts. *J Stored Prod Res* 97:101973 (2022).
- 24. Perkin LC, Friesen KS, Flinn PW, Oppert B, Venom gland components of the ectoparasitoid wasp,
 Anisopteromalus calandrae. J Venom Res 6:19–37 (2015).
- 25. Choi WI, Yoon TJ, Ryoo MI, Host-size-dependent feeding behavior and progeny sex ratio of
 Anisopteromalus calandrae (Hym., Pteromalidae). *J Appl Entomol* 125:71–77 (2001).
- Schmale I, Wäckers FL, Cardona C, Dorn S, Control potential of three hymenopteran parasitoid
 species against the bean weevil in stored beans: The effect of adult parasitoid nutrition on longevity
 and progeny production. *Biol Control* 21:134–139 (2001).
- 394 27. Menon A, Flinn PW, Dover BA, Influence of temperature on the functional response of
 395 Anisopteromalus calandrae (Hymenoptera: Pteromalidae), a parasitoid of *Rhyzopertha dominica* 396 (Coleoptera: Bostrichidae). J Stored Prod Res 38:463–469 (2002).
- 39728. Clausen CP, Introduced Parasites and Predators of Arthropod Pests and Weeds: A World Review.
- 398 United States Department of Agriculture Handbook (1978).
- 29. Van Huis A, Biological methods of bruchid control in the tropics: A review. *Insect Sci Appl* 12:87–
 400 102 (1991).
- 401 30. Tuda M, Chou L-Y, Niyomdham C, Buranapanichpan S, Tateishi Y, Ecological factors associated
- with pest status in *Callosobruchus* (Coleoptera: Bruchidae): High host specificity of non-pests to
 Cajaninae (Fabaceae). *J Stored Prod Res* 41:31–45 (2005).
 - 14

- 404 31. Tuda M, Shimada M, Complexity, evolution, and persistence in host-parasitoid experimental systems
- 405 with *Callosobruchus* beetles as the host. *Adv Ecol Res* 37:37–75 (2005).
- 406 32. Tuda M, Rönn J, Buranapanichpan S, Wasano N, Arnqvist G, Evolutionary diversification of the bean
- 407 beetle genus *Callosobruchus* (Coleoptera: Bruchidae): Traits associated with stored-product pest
 408 status. *Mol Ecol* 15:3541–3551 (2006).
- 409 33. Miksanek JR, Tuda M, Endosymbiont-mediated resistance to entomotoxic nanoparticles and sex-
- 410 specific responses in a seed beetle. PREPRINT (Version 1) available at Research Square:
- 411 <u>https://doi.org/10.21203/rs.3.rs-1941333/v1</u>
- 412 34. Helmy EAM, San PP, Zhang YZ, Adarkwah C, Tuda M, Fungus-synthesized nanoparticle and their
 413 target and nontarget effects on stored bean pest beetles and their parasitoid. PREPRINT (Version 1)
 414 available at Research Square: https://doi.org/10.21203/rs.3.rs-2097277/v1
- 415 35. El-Ramady HR, Domokos-Szabolcsy É, Abdalla NA, Alshaal TA, Shalaby TA, Sztrik A, et al.,
 416 Selenium and nano-selenium in agroecosystems. *Environ Chem Lett* 12:495–510 (2014).
- 417 36. Mechora Š, Selenium as a protective agent against pests: A review. *Plants* 8:262 (2019).
- 418 37. Garza-García JJO, Hernández-Díaz JA, Zamudio-Ojeda A, León-Morales JM, Guerrero-Guzmán A,
 419 Sánchez-Chiprés DR, et al., The role of selenium nanoparticles in agriculture and food technology.
 420 *Biol Trace Elem Res* 200:2528–2548 (2022).
- 38. Simmons TW, Jamall IS, Lockshin RA, Accumulation, distribution and toxicity of selenium in the
 adult house fly, *Musca domestica. Comp Biochem Physiol* 91: 559–563 (1988).
- 423 39. Hogan GR, Razniak HG, Selenium-induced mortality and tissue distribution studies in *Tenebrio* 424 *molitor* (Coleoptera: Tenebrionidae). *Environ Entomol* 20:790–794 (1991).
- 425 40. Lalitha K, Rani P, Narayanaswami V, Metabolic relevance of selenium in the insect *Corcyra*426 *cephalonica*: Uptake of ⁷⁵Se and subcellular distribution. Biol Trace Elem Res 41:217–233 (1994).
- 427 41. Bartůněk V, Junková J, Šuman J, Kolářová K, Rimpelová S, Ulbrich P, Sofer Z, Preparation of
- 428 amorphous antimicrobial selenium nanoparticles stabilized by odor suppressing surfactant
 429 polysorbate 20. *Mater Lett* 152:207–209 (2015).
- 430 42. Vahdati M, Tohidi Moghadam T, Synthesis and characterization of selenium nanoparticles-lysozyme
- 431 nanohybrid system with synergistic antibacterial properties. *Sci Rep* 10:510 (2020).
- 43. Cedergreen N, Ritz C, Streibig JC, Improved empirical models describing hormesis. *Environ Toxicol*433 *Chem* 24:3166–3172 (2005).
- 434 44. Ritz C, Baty F, Streibig JC, Gerhard D, Dose-response analysis using R. *PLoS One* 10:e0146021
 435 (2015).

- 436 45. Golombieski RM, Graichen DÂS, da Rocha JBT, da Silva Valente VL, da Silva Loreto ÉL, Over-
- 437 activation of the *Drosophila melanogaster hsp83* gene by selenium intoxication. *Genet Mol Biol*438 31:128–135 (2008).
- 439 46. Martin-Romero FJ, Kryukov GV, Lobanov AV, Carlson BA, Lee BJ, Gladyshev VN, et al., Selenium
 440 metabolism in *Drosophila*. *J Biol Chem* 276:29798–29804 (2001).
- 441 47. Golubkina N, Sheshnitsan S, Kapitalchuk M, Ecological importance of insects in selenium biogenic
 442 cycling. *Int J Ecol* 2014:835636 (2014).
- 443 48. Reich HJ, Hondal RJ, Why nature chose selenium. ACS Chem Biol 11:821–841 (2016).
- 444 49. Vickerman DB, Trumble JT, George GN, Pickering IJ, Nichol H, Selenium biotransformations in an
 445 insect ecosystem: Effects of insects on phytoremediation. *Environ Sci Technol* 38:3581–3586 (2004).
- 446 50. Bañuelos GS, Vickerman DB, Trumble JT, Shannon MC, Davis CD, Finley JW, et al., Biotransfer
- possibilities of selenium from plants used in phytoremediation. *Int J Phytoremediation* 4:315–329
 (2002).
- 51. Vickerman DB, Trumble JT, Biotransfer of selenium: Effects on an insect predator, *Podisus maculiventris. Ecotoxicology* 12:497–504 (2003).
- 451 52. Butler CD, Trumble JT, Predicting population dynamics of the parasitoid *Cotesia marginiventris*452 (Hymenoptera: Braconidae) resulting from novel interactions of temperature and selenium.
 453 *Biocontrol Sci Technol* 20:391–406 (2010).
- 454 53. Salem SS, Fouda MMG, Fouda A, Awad MA, Al-Olayan EM, Allam AA, et al., Antibacterial,
 455 cytotoxicity and larvicidal activity of green synthesized selenium nanoparticles using *Penicillium*456 *corylophilum. J Clust Sci* 32:351–361 (2021).
- 457 54. Wäckers FL, The effect of food supplements on parasitoid–host dynamics. *Proceedings of the First*458 *International Symposium on Biological Control of Arthropods*, Honolulu, Hawaii, pp. 226-231
- 459 (2003).
- 460 55. Miksanek JR, Heimpel GE, A matrix model describing host–parasitoid population dynamics: The
 461 case of *Aphelinus certus* and soybean aphid. *PLoS One* 14:e0218217 (2019).
- 462 56. Miksanek JR, Heimpel GE, Density-dependent lifespan and estimation of life expectancy for a
- 463 parasitoid with implications for population dynamics. *Oecologia* 194:311–320 (2020).
- 464 57. Chapple CE, Guigó R, Relaxation of selective constraints causes independent selenoprotein extinction
 465 in insect genomes. *PLoS One* 3:e2968 (2008).
- 466 58. Lobanov AV, Hatfield DL, Gladyshev V, Selenoproteinless animals: Selenophosphate synthetase
- 467 SPS1 functions in a pathway unrelated to selenocysteine biosynthesis. *Protein Sci* 17:176–182
- 468 (2008).

- 469 59. Alburaki M, Smith KD, Adamczyk J, Karim S, Interplay between Selenium, selenoprotein genes, and
- 470 oxidative stress in the honey bee *Apis mellifera* L. *J Insect Physiol* 117:103891 (2019).
- 471 60. Shchedrina VA, Kabil H, Vorbruggen G, Lee BC, Turanov AA, Hirosawa-Takamori M, et al.,
- 472 Analyses of fruit flies that do not express selenoproteins or express the mouse selenoprotein,
- 473 methionine sulfoxide reductase B1, reveal a role of selenoproteins in stress resistance. *J Biol Chem*
- 474 286: 29449 29461 (2011).
- 475 61. Santesmasses D, Mariotti M, Gladyshev VN, Bioinformatics of selenoproteins. *Antioxid Redox Signal*
- 476 33:525–536 (2020).

- 477 **Table 1.** Parameter estimates for the Cedergreen–Ritz–Streibig-modified log-logistic model for the
- 478 concentration-dependent effect of selenium nanoparticles $(0-1000 \text{ mg L}^{-1})$ on the lifespan of the
- 479 parasitoid *Anisopteromalus calandrae* in the absence of hosts.

Parameter	Grouping (sex)	Estimate	SE	<i>t</i> -value	<i>p</i> -value
b	Female	0.47	0.09	5.29	< 0.001
	Male	0.84	0.16	5.24	< 0.001
d (upper asymptote)	Female	4.55	0.15	29.60	< 0.001
	Male	3.01	0.16	19.21	< 0.001
е	_	494.80	118.95	4.16	< 0.001
f (effect of hormesis)	_	1.63	0.44	3.73	< 0.001

481 **Table 2.** Statistical tests of the effects of selenium nanoparticles (25 mg L^{-1}) on the life history

Response (Model)	Predictor	df	χ^2	<i>p</i> -value
Age at death (Cox prop. hazards)	SeNP treatment	1	4.27	0.039
Reproduction (linear regression)	SeNP treatment	1	16.24	< 0.001
Sex ratio (binomial GLM)	SeNP treatment	1	0.89	0.346
Development time (Cox prop. hazards)	Sex	1	18.38	< 0.001
	SeNP treatment	1	0.03	0.856
	Sex \times SeNP treatment	1	0.02	0.884

482 parameters of adult female parasitoids (*Anisopteromalus calandrae*) and offspring.

484 **Table 3a.** Correlation between early and late reproduction (number of offspring per day) by female

Predictor	df	χ^2	<i>p</i> -value
Early reproductive rate	1	42.97	< 0.001
[log(age-adjusted number of offspring per day)]			
SeNP treatment	1	6.32	0.016
Early reproductive rate × SeNP treatment	1	0.00	0.946

485 Anisopteromalus calandrae and the effects of selenium nanoparticles (25 mg L^{-1}).

486

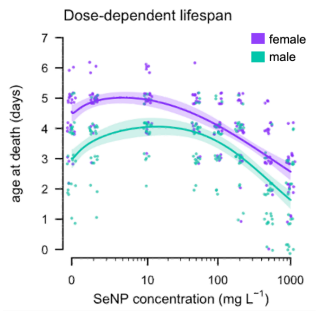
- 487 **Table 3b.** Parameter estimates for the Michaelis–Menten-type model of the lifetime reproductive success
- 488 of selenium nanoparticle-treated (25 mg L^{-1}) female parasitoids (*Anisopteromalus calandrae*).

Parameter	Estimate	SE	<i>t</i> -value	<i>p</i> -value
Maximum number of parasitized hosts, <i>m</i>	44.72	17.85	2.51	< 0.001
β	21.94	11.91	1.84	< 0.001
Preoviposition period, $t_{\rm pre}$	1.90	0.30	6.35	< 0.001
Effect of SeNP treatment, γ	0.02	0.01	3.62	< 0.001

490 **Table 4.** Statistical tests of the effects of the parasitoid *Anisopteromalus calandrae* and selenium

Response (Model)	Predictor	df	χ^2	<i>p</i> -value
Host emergence	Parasitoid presence	1	26.91	< 0.001
(LMM)	SeNP treatment	1	0.61	0.436
	Parasitoid presence × SeNP treatment	1	8.14	0.004
Host sex ratio	Parasitoid presence	1	0.06	0.808
(binomial GLMM)	SeNP treatment	1	9.22	0.002
	Parasitoid presence × SeNP treatment	1	0.13	0.715

491 nanoparticles (25 mg L^{-1}) on the host/pest (azuki bean beetle) population.



493

494 **Figure 1.** Dose-dependent lifespan of the parasitoid *Anisopteromalus calandrae* (adult lifespan, i.e. age at

death) treated individually with different concentrations of selenium nanoparticles (SeNPs) in the absence

496 of hosts; the solid line plots the Cedergreen–Ritz–Streibig-modified log-logistic model (with 95% CI);

497 points plot observed values (jittered for clarity).

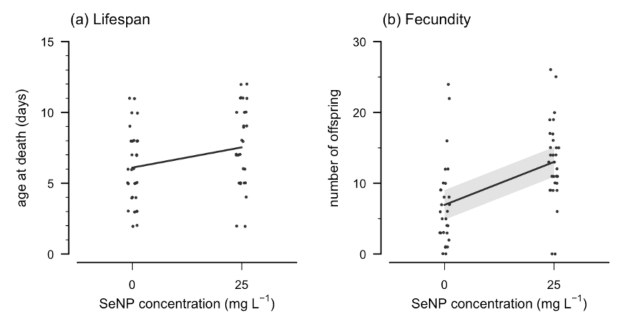


Figure 2. Lifespan and fecundity of adult female *Anisopteromalus calandrae* as a function of selenium nanoparticle (SeNP) treatment (0 or 25 mg L⁻¹). (a) Lifespan of adult female parasitoids; the solid line indicates the average age at death. (b) Fecundity (total number of offspring); the solid line plots the linear regression model (with 95% CI). In both panels, points plot observed values (jittered for clarity).

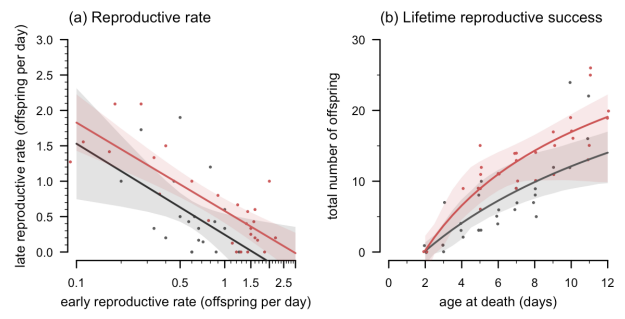
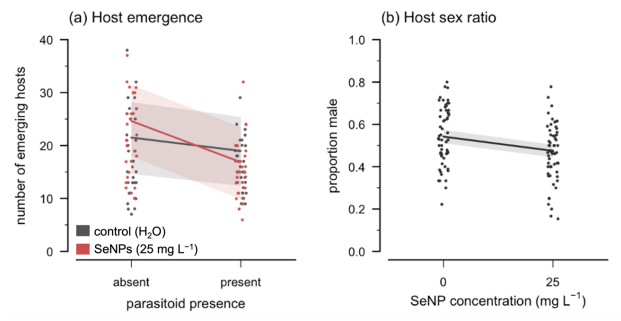


Figure 3. Reproduction of the generalist parasitoid *Anisopteromalus calandrae* as a function of age and selenium nanoparticle (SeNP) treatment (0 or 25 mg L⁻¹). **(a)** Correlation between the early and late reproductive rates of female parasitoids (number of offspring per day [age-adjusted], days 0–3 vs. day 4– death); the solid line plots the linear regression model (with 95% CI). **(b)** Lifetime reproduction, or the total number of offspring [per female]; solid lines plot the Michaelis-Menten-type models (with 95% CI). In both panels, points plot observed values (jittered for clarity in panel b).



511 **Figure 4.** Influence of parasitism and selenium nanoparticle (SeNP) treatment (0 or 25 mg L^{-1}) on the

512 host/pest population. (a) Host suppression, or the number of emerging adult azuki bean beetles; solid lines

513 plot the population-level predictions from the LMM (with 95% prediction interval). (b) Sex ratio of

514 emerging beetles; the solid line plots the population-level predictions from the binomial GLMM (with

515 95% prediction interval). In both panels, points plot observed values (jittered for clarity).