1	Pesticide-induced resurgence of the brown planthopper is mediated by
2	diverse actors that promote juvenile hormone biosynthesis and female
3	fecundity
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13 Abstract

14	Pesticide-induced resurgence, increases in pest insect populations following
15	pesticide application, is a serious threat to the sustainable control of many highly
16	damaging crop pests. Resurgence can result from pesticide-enhanced pest
17	reproduction, however, the molecular mechanisms mediating this process remain
18	unresolved. Here we show that brown planthopper (BPH) resurgence following
19	emamectin benzoate (EB) exposure results from the coordinated action of a diverse
20	suite of actors that regulate juvenile hormone (JH) levels, resulting in increased JH
21	titer in adult females and enhanced fecundity. Following reports of BPH resurgence in
22	rice crops when this species is exposed to EB, we demonstrate that EB treatment
23	results in profound changes in female BPH fitness including enhanced ovary
24	development and elevated egg production. This enhanced reproductive fitness
25	results from the EB-mediated upregulation of key genes involved in the regulation of
26	JH, including JHAMT, Met and Kr-h1 and the downregulation of allatostatin (AstA)
27	and allatostatin receptor (AstAR) expression. The remodulation of gene expression
28	following EB exposure is dependent on the action of this insecticide on its molecular
29	target the glutamate-gated chloride channel (GluCl) receptor. Collectively, these
30	results provide mechanistic insights into the regulation of negative pesticide-induced
31	responses in insects and reveal the key actors involved in the JH-signaling pathway
32	that underpin pesticide resurgence.

34 Introduction

Chemical pesticides remain the primary means of controlling many of the world's 35 36 most damaging arthropod crop pests [1, 2]. However, pesticide applications can result in pest resurgence, increases in pest insect populations that exceed natural, untreated 37 population sizes, following an initial reduction of the pest population [3-5]. Two 38 mechanisms have been implicated in pest resurgence - the loss of beneficial insects 39 including natural enemies and pesticide-enhanced pest reproduction [3]. In the case 40 of the latter, several pesticides, such as the insecticides triazophos, deltamethrin and 41 42 the fungicide jinggangmycin, have been reported to stimulate pest reproduction [6-8]. Pesticide-enhanced pest reproduction has been linked to changes in physiology and 43 biochemistry of pest organisms after exposure to pesticides [3, 4]. However, the 44 45 molecular mechanisms underlying enhanced reproduction associated with pest resurgence remain poorly resolved. 46

The brown planthopper (BPH), Nilaparvata lugens (Stål), is a notorious pest of 47 48 rice crops throughout Asia causing annual losses of ~300 million dollars across major 49 rice producing countries [2, 3]. BPH inhibits the growth of rice plants by feeding, and also transmits highly damaging plant viruses including rice grassy stunt virus and rice 50 ragged stunt virus [9]. Currently, chemical insecticides play an indispensable role in 51 52 the control of BPH due to their efficiency, rapid effect, and low cost. However, due to the widespread and intensive use of chemical insecticides, BPH has developed 53 54 resistance to the majority of compounds used for control [2, 10].

55 Emamectin benzoate (EB) and abamectin are avermectin pesticides, and act as

allosteric modulators of insect glutamate gated chloride channels (GluCls), inhibiting 56 muscle contractions that lead to the cessation of insect feeding and subsequent death 57 58 [11]. These insecticides exhibit particularly strong activity against Lepidoptera such as the rice leaffolder, Cnaphalocrocis medinalis Guénee, an important foliage-feeding 59 insect which attacks rice during the vegetative stage [12]. Both BPH and the rice 60 leaffolder are migratory pests with overlapping migratory paths, however, their 61 occurrence period in the field differs by approximately one month, with leaffolders 62 appearing earlier than BPH. Therefore, the use of EB to control rice leaffolder has the 63 64 potential to impact BPH arriving later, via exposure to sublethal concentrations of this compound. In this regard, we have observed that when farmers use EB and abamectin 65 to control leaffolders on rice crops in China, BPH outbreaks frequently occur in the 66 67 same field. While sublethal doses of certain pesticides have been shown to enhance fecundity in BPH, including the insecticides triazophos and deltamethrin [6, 7, 13, 14] 68 and the fungicides carbendazim and jinggangmycin [8], whether avermectins trigger 69 70 resurgence in BPH via insecticide-enhanced reproduction remains unclear.

Reproduction in insects is influenced by external factors such as light [15], temperature [16], humidity [17] and nutrition [18], and endogenous factors such as the juvenile hormone (JH) [19], ecdysone [20], insulin [21] and TOR [22] pathways [23, 24]. Of these, JH, has been particularly implicated in insecticide-induced enhanced fecundity, with triazophos and deltamethrin treatments leading to increased circulating JH III titers in BPH females [3]. JH is synthesized and secreted by the corpora allata in insects [25], and can promote reproduction by regulating the synthesis and secretion of vitellogenin (vg) in the female fat body, and stimulating the absorption of vg by the
developing oocyte [19]. However, the regulation of JH is complex [19] and the key
actors involved in JH-mediated pesticide-enhanced reproduction remains an open
question.
In this study, we used a diverse range of approaches to investigate the impact of

sublethal doses of avermectins on BPH fecundity, and unravel the molecular mechanisms mediating enhanced reproduction following exposure to this insecticide class. We show that avermectin exposure results in profound changes in the expression of a key suite of genes that in combination regulate JH, resulting in increased JH titer in adult females, which promotes fecundity.

89 RESULTS

90 GluCl allosteric modulators (emamectin benzoate and abamectin) stimulate

91 fecundity of female *N. lugens*

92	To investigate whether GluCl modulators affect fecundity in BPH, we first
93	determined the sub-lethal and median lethal doses of emamectin benzoate (EB) to
94	4 th instar nymphs, newly emerged males and females of BPH (Table S1). For this we
95	employed two different bioassay methods, the rice seedling dip bioassay method and
96	topical application bioassay method [2, 26], in order to assess both the systemic and
97	contact toxicity of these insecticides (Table S1). We then systemically treated 4^{th}
98	instar nymphs of BPH with the estimated LC_{15} and LC_{50} concentrations of EB and
99	examined the fecundity of BPH after these individuals mated with treated or
100	untreated individuals. We use the term "t" to represent individuals treated with EB
101	and "ck" to indicate control individuals that were treated with insecticide diluent minus
102	insecticide. After treatment with the LC_{15} and LC_{50} concentrations of EB the number
103	of eggs laid per female of BPH in \bigcirc t × \eth t crosses increased by 1.48 and 1.40 times
104	compared with control \bigcirc ck × \bigcirc ck crosses (Figure 1A); the number of eggs laid per
105	female of BPH in \bigcirc t × \bigcirc ck crosses increased by 1.53 and 2.07 times compared with
106	control crosses (Figure 1B); However, the number of eggs laid by per female of BPH
107	in \bigcirc ck × \bigcirc t crosses did not increase significantly compared to control \bigcirc ck × \bigcirc ck
108	crosses (Figure 1C).

		LC ₁₅ (95%F.L.)	LC ₅₀ (95%F.L.)		
Development	Slope ± SE	(mg/L) or LD ₁₅	(mg/L) or LD ₅₀	. 2(-15)	Duratura
al stages		(95%F.L.)	(95%F.L.)	χ²(at)	P value
		(pg/insect)	(pg/insect)		
4 th nymph	0.50 + 0.407	0.87	2.21 (1.53-	0.00 (4)	0.04
	2.56 ± 0.487	(0.39-1.32)	2.82)	2.68 (4)	0.61
Male	2.42 ± 0.35	0.80	2.13 (1.48-	2.12 (4)	0.72
		(0.40-1.20)	2.78)		
Female 2.4		2.33	6.12 (4.18-	3.56 (4)	0.50
	2.47 ± 0.41	(1.09-3.56)	8.00)		
4 th nymph 2.52 ± 0.35	0.50 . 0.05	15.54 (9.46-	39.90 (30.66-		0.63
	2.52 ± 0.35	21.42)	49.56)	2.60 (4)	
Male 2.77 ± 0.57	6.72	15.96	4.00 (0)	0.75	
	2.//±0.5/	(2.52-10.92)	(9.66-21.84)	1.23 (3)	0.75
	Development al stages 4 th nymph Male Female 4 th nymph Male	Development al stages $\operatorname{Slope \pm SE}$ 4^{th} nymph 2.56 ± 0.487 Male 2.42 ± 0.35 Female 2.47 ± 0.41 4^{th} nymph 2.52 ± 0.35 Male 2.77 ± 0.57	$\begin{array}{c} \mbox{LC}_{15} \ (95\% F.L.) \\ \mbox{Implement} \\ a \ stages \end{array} & \begin{array}{c} \mbox{Implement} \\ \mbox{Slope \pm SE} \\ a \ stages \end{array} & \begin{array}{c} \mbox{Implement} \\ \mbox{(mg/L) or LD}_{15} \\ \mbox{(95\% F.L.)} \\ \mbox{(pg/insect)} \end{array} \\ \mbox{Implement} \end{array} & \begin{array}{c} \mbox{Implement} \\ \mbox{(pg/insect)} \end{array} \\ \mbox{Implement} \end{array} & \begin{array}{c} \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} & \begin{array}{c} \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} & \begin{array}{c} \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} & \begin{array}{c} \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} & \begin{array}{c} \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} & \begin{array}{c} \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} & \begin{array}{c} \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} & \begin{array}{c} \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} \\ \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} \\ \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} \\ \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{aligned} \\ \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \end{array} \\ \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \\ \mbox{Implement} \mbox{Implement} \\ $	$\begin{array}{c} \mbox{LC}_{15} \ (95\% F.L.) & \mbox{LC}_{50} \ (95\% F.L.) \\ \mbox{Imm} \mbox{Imm} \ (100 \ 100 $	$ \begin{array}{c} \label{eq:constraints} \\ \mbox{Development} \\ \mbox{al stages} \end{array} & \begin{array}{c} \mbox{LC}_{15} (95\% F.L.) & \mbox{LC}_{50} (95\% F.L.) & \mbox{(mg/L) or LD}_{50} \\ \mbox{(mg/L) or LD}_{15} & \mbox{(mg/L) or LD}_{50} \\ \mbox{(95\% F.L.)} & \mbox{(95\% F.L.)} & \mbox{(95\% F.L.)} & \mbox{(pg/insect)} & \mbox$

Table S1. Determination of the toxicity of emamectin benzoate against BPH in systemic and topical application bioassays.

110			40.40.(0.70	27.00		
111	Female	3.31 ± 0.76	18.48 (6.72-	37.80	0.79 (3)	0.85
112			20.14)	(23.10-30.40)		

113	Exposure of 4^{th} instar nymphs with the LC ₁₅ and LC ₅₀ concentrations of EB in
114	contact bioassays also significantly stimulated fecundity. After treatment with the LC_{15}
115	and LC_{50} concentrations of EB, the number of eggs laid by per female of BPH in ${\mathbb Q}t$ ×
116	\exists t crosses increased by 1.18 and 1.26 times compared with the control (\bigcirc ck × \exists ck)
117	(Figure 1D); The number of eggs laid per female of BPH in $\operatorname{Pt} \times \operatorname{Cck}$ crosses
118	increased by 1.27 and 1.56 times compared with the control crosses (Figure 1E);
119	However, there was no significant difference in egg-laying number between \Im ck × \Im t
120	crosses and controls (♀ck × ♂ck) (Figure 1F). These results reveal that EB can
121	stimulate the fecundity of females following both the systemic and contact routes of
122	exposure.





Figure 1. Fecundity of BPH following exposure to sub-lethal (LC₁₅) and median lethal (LC₅₀)
concentrations of emamectin benzoate following system application bioassays (A: ♀t × \$t;
B: ♀t × \$ck; C: ♀ck × \$t\$) and topical application bioassays (D: ♀t × \$t; E: ♀t × \$ck; F:

- 128 \bigcirc ck × \Diamond t), respectively. The letter "t" represents treatment with insecticide, while "ck"
- 129 indicates controls that was not treated with insecticide. All data are presented as the mean ±
- 130 s.e.m. Different lower-case letters above the bars indicate significant differences (One-way
- 131 ANOVA with Tukey's Multiple Range Test, p < 0.05).
- 132

141

- 133 We next examined whether EB treatment of adult BPH also stimulates
- reproduction. Indeed, treating newly emerged adults with the LC₁₅ and LC₅₀
- concentrations of EB significantly stimulated the number of eggs laid per female
- 136 (Figure 1-figure supplement 1A). Furthermore, sub-lethal exposure of 4th instar BPH
- 137 nymphs to another GluCl allosteric modulator, abamectin (LC₁₅ and LC₅₀
- 138 concentrations) was also found to significantly enhance reproduction (Figure 1-figure
- 139 supplement 1B).
- 140 Figure 1-figure supplement 1



Figure 1-figure supplement 1. (A) Fecundity of BPH when newly emerged adults were

treated with sub-lethal (LD₁₅) and median lethal (LD₅₀) concentrations of emamectin benzoate

144 via topical application. (B) Fecundity of BPH when 4th instar nymphs were treated with sublethal (LC₁₅) and median lethal (LC₅₀) concentrations of abamectin via systemic exposure. All 145 146 data are presented as the mean ± s.e.m. Different lower-case letters above the bars indicate significant differences (One-way ANOVA with Tukey's Multiple Range Test, p < 0.05). 147 To examine if EB also stimulates egg-laying in other insect species we 148 conducted bioassays on the small brown planthopper, Laodelphax striatellus, the 149 white backed planthopper, Sogatella furcifera and fruit flies Drosophila melanogaster. 150 In contrast to our findings on BPH, we found that sub-lethal doses (LC_{15} and LC_{50}) of 151 EB inhibits fecundity of female *L. striatellus*, (Figure 1-figure supplement 2A-C) and 152 153 has no impact on the fecundity of S. furcifera, (Figure 1-figure supplement 2D-F). In addition, we found that sublethal doses (LC_{15} or LC_{50}) of EB also inhibit fecundity in 154 D. melanogaster (Figure 1-figure supplement 2G and H). These results indicate that 155 156 the stimulation of reproduction by EB in BPH is species-specific and does not extend to even related insect species. 157

158 Figure 1-figure supplement 2



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160 **Figure 1-figure supplement 2.** Fecundity of small brown planthopper, *Laodelphax striatellus*,

161 (A-C) white backed planthopper, Sogatella furcifera (D-F) and fruit fly, Drosophila

162 *melanogaster* (G and H) when larvae and newly emerged adults were treated with sub-lethal

- 163 concentrations of emamectin benzoate. All data are presented as the mean ± s.e.m. Different
- 164 lower-case letters above the bars indicate significant differences (One-way ANOVA with
- 165 Tukey's Multiple Range Test, p < 0.05).



168	To better understand the effects of EB on the reproductive fitness of BPH, the
169	preoviposition period, emergence rate, female ratio, female longevity and female
170	weight were evaluated following exposure using the systemic bioassay. The
171	preoviposition period of females treated with the LC_{50} of EB decreased significantly
172	compared with the control (Figure 1-figure supplement 3A). In contrast no significant
173	effects of EB on emergence rate and female ratio were observed (Figure 1-figure
174	supplement 3B and C). Female survival analysis showed that exposure of 4 th instar
175	nymphs to the LC_{50} of EB has no impact on female longevity (Figure 1-figure
176	supplement 3D). Interestingly, brachypterism (long-wing) female ratio and female
177	weight were significantly increased after EB exposure (Figure 1-figure supplement 3E
178	and F).



181

180

Figure 1-figure supplement 3. The impact of emamectin benzoate on the reproductive 182

fitness of BPH. Fourth instar nymphs were treated with the LC₅₀ concentration of emamectin 183

184 benzoate in systemic bioassays. All data are presented as the mean ± s.e.m. Different lower-

185 case letters above the bars indicate significant differences (Student's *t* test, p < 0.05).

186

EB promotes ovarian development in BPH 187

Figure 1-figure supplement 3

188	To investigate the cause of increased female weight following EB exposure we
189	examined if EB influences ovary development in BPH. We dissected and compared
190	the ovaries of females treated with the LC_{50} of EB at 1, 3, 5 and 7 days after eclosion
191	(DAE) with control females. At 3, 5 and 7 DAE, the number of detained eggs of BPH

- in the EB treated group were significantly higher than that of controls (Figure 2A and
- B). We also explored whether EB treatment could enhance or impair oogenesis in
- BPH. However, dissection of various developmental stages revealed that emamectin
- 195 benzoate treatment has no significant effects on oogenesis (Figure 2C and D).

196 Figure 2



197

Figure 2. The impact of emamectin benzoate on ovary development in BPH. Fourth instar 198 nymphs were treated with the LC₅₀ concentration of emamectin benzoate in systemic bioassays. 199 200 (A) Ovary development in EB treated BPH at 1, 3, 5 and 7 days after eclosion (DAE) compared 201 to untreated controls. Scale bar = 1,000 µm. (B) Number of detained eggs in the ovaries of EB treated BPH females measured at 1, 3, 5 and 7 DAE compared to controls. All data are 202 203 presented as the mean ± s.e.m. Asterisks indicate values significantly different from the control using student *t* test (ns, no significant; *p < 0.05 and **p < 0.01). (C) Different developmental 204 205 stages of BPH eggs. (D) No impairment of emamectin benzoate on oogenesis of BPH. Scale 206 bar = 100 µm.

208 EB exposure enhances the abundances of storage macromolecules and

209 circulating sugars in BPH

Nutritional status is an important indicator of reproductive fitness. Thus, to investigate whether EB affects intermediary metabolism and energy storage of BPH, glycogen, triacylglyceride (TAG), total protein content, cholesterol and four circulating carbohydrates were quantified in 4th instar BPH nymphs following exposure to the LC₅₀ of EB.

215	We found that EB exp	sure has no impa	ct on glycogen leve	ls (Figure 2-figure
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supplement 1A). The amounts of TAG in EB-treated BPH were 27% higher (p < 0.05)

than those in controls, but only in BPH of the late fifth instar (5L) stage, with no

significant differences observed in subsequent developmental stages (Figure 2-figure

supplement 1B). The amount of total protein content in EB-treated BPH was higher

than the control groups in the case of all developmental stages from 5L nymph to

221 7DAE (Figure 2-figure supplement 1C). EB exposure also increased cholesterol

levels at 4 and 5 DAE (Figure 2-figure supplement 1D). Compared with the solvent

223 control, EB treatment caused significant increases (p < 0.05) in the levels of sucrose,

glucose, fructose, and trehalose (Figure 2-figure supplement 1E-H). Thus,

collectively, these data provide evidence that EB exposure leads to energy

226 mobilization and the metabolism of carbohydrates and lipids in BPH.



228 Figure 2-figure supplement 1

Figure 2-figure supplement 1. Amounts of Glycogen (A), TAG (B), total protein content (C), cholesterol (D) and four circulating sugars including sucrose, glucose, fructose and trehalose (E-H) after BPH exposure to EB. All data are presented as the mean \pm s.e.m. The differences between the EB-treated and solvent-treated BPH were analyzed using unpaired student *t*-test (*, *p* < 0.05; **, *p* < 0.01; ***, *p* < 0.001; ****, *p* < 0.0001).

235

236 **EB stimulates egg-laying that is mediated by the JH signaling pathway**

237	Given the important role of juvenile hormone (JH) in vitellogenesis and egg
238	development in insects [19, 25, 27-29], we asked whether EB-treatment could
239	influence the titer of JH in BPH. As measured by ELISA, the juvenile hormone titer of
240	BPH nymphs treated with the LC_{50} concentration of EB was significantly lower than
241	that of controls in systemic bioassays during the middle and late stages of the 4^{th}
242	instar (Figure 3A). However, at 2, 3 and 4 DAE, the JH titer in the EB treated group
243	was significantly higher than that of the control (Figure 3A). Interestingly, the titer of
244	another important insect hormone, the steroid ecdysone, was not significantly
245	different between EB-treated BPH and solvent-treated controls (Figure 3-figure

246	supplement 1). To independently validate the results of ELISA assays, we employed
247	HPLC-MS/MS to measure JH titer in BPH following EB exposure [27, 30, 31]. The
248	results showed that JH III titer significantly decreased after EB-treatment at the late
249	4 th instar nymph stage (Figure 3A and B), but significantly increased at the third day
250	after eclosion (3 DAE) (Figure 3A and C). To further investigate the role of JH in EB-
251	enhanced fecundity in BPH, we treated BPH with methoprene and pyriproxyfen, JH
252	analogues or biologically active JH mimics respectively, to determine whether they
253	can stimulate fecundity in BPH. Both compounds significantly increased egg-laying in
254	BPH (Figure 3D). Takes together these results reveal that EB stimulates an
255	increased in JH titer that induces fecundity in BPH. Since we found that EB could
256	induce JH synthesis in the BPH, we asked whether EB could influence the
257	expression of genes that are involved in JH synthesis or degradation. For this we
258	treated 4^{th} instar nymphs of BPH with the LC ₅₀ concentration of EB using systemic
259	bioassays, and then collected early (5E), middle (5M) and late (5L) stage of 5^{th} instar
260	nymph and 1-7 DAE female adults (DAE) for analysis. Quantitative PCR was then
261	used to examine the expression of key genes previously implicated in the regulation
262	of JH [<u>32</u> , <u>33</u>].
263	Farnesoic acid O-methyltransferase (FAMeT) [<u>34</u>], is known as an important
264	enzyme in the JH biosynthetic pathway, catalyzing methylation of farnesoic acid (FA)
265	to methyl farnesoate (MF) [33]. We found that this gene was significantly upregulated

in 5M instar nymph to 4 DAE after EB- treatment (1.0-fold to 3.0-fold) (Figure 3E). 266

Another gene, juvenile hormone acid methyltransferase (JHAMT) [33, 35], which is 267

268	involved in the biosynthesis of JH, was also upregulated in EB-treated BPH at 1 to 5
269	DAE than controls (1.5-fold to 3.0-fold) (Figure 3F). Methoprene-tolerant (Met),
270	belongs the basic helix-loop-helix Per/Arnt/Sim (bHLH-PAS) family of transcription
271	factors and is the intracellular (nuclear) receptor of JH [<u>36</u> , <u>37</u>]. The levels of <i>met</i>
272	mRNA increased in EB-treated BPH at the 5M and 5L instar nymph and 1 to 5 DAE
273	stages compared to controls (1.7-fold to 2.9-fold) (Figure 3G). Krüppel homolog 1
274	(Kr-h1), a transcriptional target of JH signaling, is reported to be sensitive to levels of
275	JH and its expression levels are directly correlated with JH titers [<u>32</u> , <u>38</u> , <u>39</u>]. We
276	found that Kr-h1 was significantly upregulated in the adults of EB-treated BPH at the
277	5M, 5L nymph and 4 to 5 DAE stages (4.7-fold to 27.2-fold) (Figure 3H). Similarly, the
278	expression levels of vitellogenin (Vg), a key downstream component of JH signaling
279	triggering ovary development in insects including BPH [40], was markedly increased
280	in females at 2–5 DAE by EB (1.7-fold to 5.5-fold) (Figure 3I). Juvenile hormone
281	esterase (JHE) is the primary JH-specific degradation enzyme that plays a key role in
282	regulating JH titers [41]. Interestingly, we observed a significant upregulation of JHE
283	mRNA levels in the early and middle 5 th instar nymph stage followed by
284	downregulation in 5L instar nymphs of EB-treated BPH (1.3-fold to 2.6-fold) (Figure
285	3J). In combination these results reveal that EB has profound impacts on the
286	expression of key genes involved in the synthesis of JH or downstream signaling
287	pathway genes that might promote egg development.
288	To further understand whether these JH pathway-related genes were involved in
289	egg-laying behavior in BPH, we performed RNAi experiments to downregulate the

expression of JHAMT, Met and Kr-h1 (Figure 3-figure supplement 2A-C). We found

- that silencing of these three genes downregulated the titer of JH in BPH providing
- functional evidence of their role in the regulation of JH (Figure 3-figure supplement
- 20). Furthermore, silencing of *Kr-h1* inhibited *FAMeT* and *Vg* gene expression while
- increasing *JHE* gene expression (Figure 3-figure supplement 2E-H). Importantly,
- silencing of JHAMT, Met and Kr-h1 gene in female BPH was also found to suppress
- egg-laying (Figure 3K-M). However, this phenotype was rescued by EB treatment,
- which restored egg-laying to normal levels in BPH injected with JHAMT, Met and Kr-
- *h1* dsRNA (Figure 3K-M). Together these results provide a mechanistic
- understanding of how EB enhances fecundity in BPH by modulating the expression
- 300 of key genes involved in JH synthesis (Figure 3N).



303

Figure 3. EB induced reproduction in BPH is mediated by components of the JH signaling pathway. (A) The titer of JH III (as measured by ELISA assay) at different developmental stages of BPH when 4th instar nymphs were treated with the median lethal concentrations of EB. (B and C) The titer of JH III (as measured by HPLC-MS/MS) in BPH females at 4L and 3 DAE when treated with median lethal concentrations of EB. (D) Oviposition rate of BPH when 4th instar nymphs were treated with 4 ppm methoprene or 10 ppm

310 pyriproxyfen. (E-J) Expression of selected JH-related genes (FAMeT, JHAMT, Met, Kr-h1, Vq, and JHE) in EB-treated BPH. (K) Egg production following silencing of JHAMT with or without 311 312 EB application. (L) Egg production following silencing of *met* with or without EB application. (M) Egg production after silencing Kr-h1 with or without EB application. (N) Schematic illustrating 313 the proposed impact of EB on the JH signaling pathway leading to enhanced reproduction. The 314 question mark indicates one or more potential additional signals. All data are presented as 315 316 means ± s.e.m. Student's t test was used to compare the two samples. One-way ANOVA with 317 Tukey's multiple comparisons test was used to compare more than two samples. ns, no significant difference; Asterisks indicate values significantly different from the control (ns, no 318 significant; *p < 0.05, **p < 0.01, ***p < 0.001 and ****p < 0.0001). Different lower-case letters 319 above the bars indicate significant differences (p < 0.05). 320

321 Figure 3-figure supplement 1





322

Figure 3-figure supplement 1. Ecdysone titer at different developmental stages of BPH when

324 4th instar nymphs were treated with median lethal concentrations of EB.

325

327 Figure 3-figure supplement 2



328

Figure 3-figure supplement 2. (A-C) Expression of JHAMT (A), Met (B) and Kr-h1 (C) following RNAi knockdown. (D) The titer of JHIII (as measured by HPLC-MS/MS) at 3 DAE when female adults were injected with dsJHAMT, dsMet, and dsKr-h1 at 1 DAE. (E-H) Expression of FAMeT, Vg, JHE and JHAMT when Kr-h1 was silenced in BPH. All data are presented as means \pm s.e.m. *p < 0.05; Mann–Whitney test.

EB induces JH biosynthesis through the peptidergic AstA/AstAR signaling pathway

- The timing and level of JH biosynthesis can be precisely regulated by the
- neuropeptides, stimulatory allatotropins (ATs) and inhibitory allatostatins (Asts), in
- many insects [42-49]. Insects can, in a species-specific manner, produce one type of
- AT and three types of Asts: FGL-amide Ast (AstA) [43, 50]; myoinhibitory peptide
- 341 (MIP or AstB) and PISCF Ast (AstC) [47, 51]. In some species, there also exists two
- paralogue genes of AstCs which are named AstCC and AstCCC [52, 53].
- Interestingly, the allatostatic activity of these three types of Ast peptides varies
- 344 between insect species so that in each species only one type of Ast (for example
- 345 AstA) controls JH production [<u>47</u>, <u>48</u>, <u>51</u>, <u>54</u>].
- Analysis of the BPH neural transcriptome sequence data has revealed the
- 347 presence of one AT, four types of Asts and four corresponding receptors, allatotropin
- receptor (A16, ATR), AstA receptor (A2, AstAR), AstB (MIP) receptor (A10, AstBR or
- MIPR) and AstC receptor (A1, AstCR) [55]. We cloned the five neuropeptide genes
- 350 (AT, AstA, AstB/MIP, AstCC and AstCCC) and confirmed the sequence obtained from
- transcriptome data (Figure 4-figure supplement 1) [55]. Interestingly, we found that
- AstC is missing in the genome of BPH and only AstCC and AstCCC are present
- 353 (Figure 4-figure supplement 1). Next, we also cloned their corresponding receptors
- ³⁵⁴ [52] including ATR (A16), AstAR (A2), AstBR (A10) and AstCR (A1) which might be
- activated by AstCC and/or AstCCC [<u>39</u>, <u>52</u>, <u>56</u>]. Sequence alignments and
- 356 phylogenetic analysis are shown in Figure 4-figure supplement 2.

357	Quantitative PCR was then used to examine if EB treatment could influence the
358	expression of the genes encoding these neuropeptides and their receptors. Treating
359	BPH with the LC_{50} concentration of EB significantly increased the expression of AT,
360	ATR and AstCCC while resulting in the downregulation of AstA, AstB/mip, AstCC,
361	AstAR and AstBR/mipr at the adult stage (Figure 4A and B, Figure 4-figure
362	supplement 3A-G). Among these, AstA and AstAR were the most downregulated
363	genes after EB treatment (Figure 4A and B, Figure 4-figure supplement 3H) and thus
364	the AstA/AstAR signaling system was selected for subsequent functional analysis.
365	Silencing of AstAR in female BPH using RNAi (Figure 4C), resulted in an increased
366	number of eggs laid per female compared with <i>dsgfp</i> -injected controls (Figure 4D).
367	Interestingly, silencing AstAR also resulted in the upregulation of FAMeT, JHAMT,
368	<i>Met</i> and <i>Kr-h1</i> which are involved in the JH biosynthesis/signaling (Figure 4E-H).
369	However, JHE was not influenced by AstAR silencing (Figure 4I). We therefore
370	investigated whether silencing the AstAR gene could influence JH titer in BPH. A
371	significantly increased JH titer was observed in AstAR silenced BPH compared with
372	controls (Figure 4J). Thus, our data strongly suggest that AstA is a key inhibitor of JH
373	production in BPH.
374	Finally, we investigated whether injection of mature Ast and AT peptides could
375	influence the number of eggs laid and JH titer in BPH. We synthesized one AT, six
376	AstAs (AstA1 to AstA6), one AstCC and one AstCCC peptide according to their
377	determined sequences (Figure 4-figure supplement 1). Indeed, we found that
378	injection of AstA1 and AstA6 reduced the number of eggs laid per female in 48 h

- 379 (Figure 4K). We also observed that AstA1 injection after 16h and 48h decreased the
- 380 JH titer significantly (Figure 4L), and AT injection increased the JH titer after 2h but
- 381 levels returned to normal after 4h injection (Figure 4M). Collectively, these results
- provide compelling evidence that EB induces reproduction in BPH through the
- 383 AstA/AstAR and JH signaling pathways (Figure 4N) and further supports the role of
- 384 AstA and AT in regulation of JH titer.

385 Figure 4

386



Figure 4. EB induced reproduction in BPH is mediated by the AstA/AstAR and JH
 signaling pathway.

(A and B) Expression of AstA and AstAR in different stages of BPH following EB treatment. (C) 389 390 Downregulation of AstAR using RNAi leads to a reduction in mRNA expression level. (D) Egg 391 production in female BPH following silencing of AstAR gene. (E-I) Expression of selected JH 392 signaling pathway related genes (FAMeT, JHAMT, Met, Kr-h1 and JHE) in AstAR silenced BPH. 393 (J) JHIII titer of BPH females after AstAR gene silencing determined by HPLC-MS/MS. (K) 394 Number of eggs laid per female in 48h following injection of six AstA1-AstA6 mature peptides 395 and one AT mature peptide. Fifty nanoliter of PBS (as control) and seven peptides (20 396 pmol/insect) were injected into female BPH three days after eclosion. (L and M) The JH III titer

397	of BPH females at different time points following AstA or AT injection. (N) Schematic of the
398	proposed role of AstA/AstAR in the regulation of JH following EB exposure. The question
399	mark indicates one or more possible additional signals. All data are presented as means \pm s.e.m.
400	Student's <i>t</i> test was used to compare the two samples. One-way ANOVA with Tukey's multiple
401	comparisons test was used to compare more than two samples. ns, no significant difference;
402	Asterisks indicate values significantly different from the control (ns, no significant; * p < 0.05, ** p
403	< 0.01, *** p < 0.001 and **** p < 0.0001). Different lower-case letters above the bars indicate
404	significant differences ($p < 0.05$).
405	

406 Figure 4-figure supplement 1



407

408 Figure 4-figure supplement 1. Alignments of the amino acid sequences of: (A) AT, (B) AstA, 409 (C) AstB/MIP, (D) AstCC and (E) AstCCC peptides from select species. AT, AstA, AstB/MIP and 410 AstCCC are predicted to have a C-terminal amide. The mature peptides belonging to the same species have been highlighted with the same color. Species names are as follows: Nillu 411 (Nilaparvata lugens), Locmi (Locusta migratoria), Scham (Schistocerca americana), Homvi 412 (Homalodisca vitripennis), Manse (Manduca sexta), Spofr (Spodoptera frugiperda), Drome 413 (Drosophila melanogaster), Spoex (Spodoptera exigua), Nasvi (Nasonia vitripennis), Grybi 414 (Gryllus bimaculatus), Bommi (Bombyx mori); Mesma (Mesobuthus martensii), Stear 415

- 416 (Stegodyphus araneomorph), Limpo (Limulus polyphemus), Carma (Carcinus maenas), Strma
- 417 (Strigamia maritima), Athro (Athalia rosae), Apime (Apis mellifera), Dapma (Daphnia magna).
- 418 Black lines under the sequences indicate the locations of the disulfide bridges in the mature
- 419 peptides. The accession numbers of the sequences are listed in Figure 4-figure supplement 1
- 420 source data.
- 421

422 Figure 4-figure supplement 2



423

Figure 4-figure supplement 2. Phylogenetic tree of the predicted BPH (*) allatotropin receptor (A16, ATR), allatostatins A receptor (A2, AstAR), AstB (MIP) receptor (A10, AstBR or MIPR) and allatostatins C receptor (A1, AstCR) with other insect species. The tree was generated using the maximum likelihood method. *Drosophila melanogaster* metabotropic glutamate receptor was included as an outgroup. The accession numbers of the sequences used for this phylogenetic tree are listed in Figure 4-figure supplement 2 source data.

430 Figure 4-figure supplement 3



431

432 Figure 4-figure supplement 3. EB induced changes in the expression of AT, AstB, AstCC,

433 **AstCCC, ATR, AstBR and AstCR in BPH.** All data are presented as means ± s.e.m. Student's

t test was used to compare the two samples. ns, no significant difference; Asterisks indicate

435 values significantly different from the control (*p < 0.05, **p < 0.01, and ***p < 0.001).

EB-enhanced fecundity in BPH is dependent on its molecular target protein the GluCl

439	EB and abamectin, are allosteric modulators, which target glutamate-gated
440	chloride channels (GluCls) [57-59]. Hence, we examined whether EB-stimulated
441	fecundity in BPH is influenced by its molecular target GluCI. The full length GluCI
442	coding sequence from BPH was cloned and sequenced (Figure 5-figure supplement
443	1) and the impact of EB on <i>GluCl</i> gene expression examined using quantitative PCR.
444	Treatment of BPH with the LC_{50} concentration of EB significantly downregulated
445	<i>GluCl</i> gene expression at the 5E and 5M nymph stages while upregulating <i>GluCl</i>
446	gene expression at 2 DAE and 5 DAE in the adult stage (Figure 5A). To examine the
447	role of GluCl gene in BPH fecundity, RNAi was used to knockdown expression of this
448	gene in female BPH (Figure 5B). A significant decrease in the number of eggs laid by
449	per female was observed in <i>dsGluCl</i> -injected insects compared with <i>dsgfp</i> -injected
450	insects (Figure 5C). However, treatment with EB was found to rescue the decreased
451	egg-laying phenotype induced by <i>dsGluCl</i> injection (Figure 5C). To investigate the
452	mechanism by which GluCl expression modulates fecundity we examined if silencing
453	GluCl influences JH titer and JH-related gene expression. Indeed, we observed that
454	RNAi knockdown of <i>GluCl</i> leads to a decrease in JH titer (Figure 5D) and down-
455	regulation of genes including JHAMT which is responsible for JH synthesis, and the
456	JH signaling downstream genes <i>Met and Kr-h1</i> (Figure 5E-G). In contrast,
457	expression of FAMeT and JHE were not changed in the GluCl silencing insects
458	(Figure 5H and I). We also examined whether silencing GluCl impacts the

- 459 AstA/AstAR signaling pathway. Silencing GluCl was found to have no impact on the
- 460 expression of *AT*, *AstA*, *AstB*, *AstCC*, *AstAR*, and *AstBR*. However, the expression of
- 461 AstCCC and AstCR was significantly upregulated in dsGluCl-injected insects (Figure
- 462 5-figure supplement 2A-H). These results suggest that EB activates *GluCl* which
- 463 induces JH biosynthesis and release, which in turn stimulates reproduction in BPH
- 464 (Figure 5J).

466 Figure 5



467

Figure 5. EB induced reproduction in brown planthoppers through its molecular target protein GluCl.

(A) Expression of GluCl in EB-treated and untreated BPH. (B) Expression of GluCl following 470 471 injection of dsGluCl in BPH. (C) Egg production after GluCl gene knockdown in EB-treated and 472 untreated BPH. (D) The JH III titer of BPH females after GluCl gene silencing as quantified using the ELISA method. (E-I) Expression patterns of selected JH-related genes (JHAMT, Met, 473 474 Kr-h1, FAMeT and JHE) in GluCl silenced BPH. (J) Schematic of the proposed role of GluCl as 475 a molecular target of EB and EB-enhanced reproduction in BPH. The question mark indicates 476 one or more possible additional signals. All data are presented as means ± s.e.m. Student's t 477 test was used to compare the two samples. One-way ANOVA with Tukey's multiple comparisons test was used to compare more than two samples. ns, no significant difference; 478 479 Asterisks indicate values significantly different from the control (ns, no significant; *p < 0.05 and **p < 0.01). Different lower-case letters above the bars indicate significant differences (p < 0.05). 480

481 Figure 5-figure supplement 1

482



Figure 5-figure supplement 1. Phylogenetic analysis of glutamate-gated chloride channels in different species. The numbers at the nodes of the branches represent the percentage bootstrap support (1000 replications) for each branch. The *Sogatella furcifera* GABA-gated chloride channel and *Nilaparvata lugens* nAchR were used as outgroup. Alignment was performed with amino acid sequences from TM1-7. The receptor names are listed in the tree. The accession numbers of the sequences used for this phylogenetic tree are listed in Figure 5figure supplement 1 source data.



Figure 5-figure supplement 2. The expression of *AT*, *AstA*, *AstB*, *AstCC*, *AstCCC*, *AstAR*, *AstBR* and *AstCR* in BPH injected with *dsGluCl* or *dsgfp*. All data are presented as means \pm s.e.m. Student's *t* test was used to compare the two samples. ns, no significant. Different lowercase letters above the bars indicate significant differences (p < 0.05).

498 Discussion

499	Pesticide-induced resurgence of pest insects is a serious problem in rice and
500	several other cropping systems [3]. However, the mechanisms underpinning pesticide-
501	enhanced reproduction in insects remain poorly understood. Here we reveal that a
502	suite of molecular actors underlie this trait that, in combination, mediate profound
503	physiological changes in the reproductive fitness of female BPH. Our data provide
504	fundamental insights into the molecular mechanisms by which xenobiotics modify
505	insect reproduction and have applied implications for the control of a highly damaging
506	crop pest. We discuss these topics below.
507	

Sublethal doses of GluCl modulators, EB and abamectin, stimulates fecundity in
 BPH

We show that in both contact and systemic assays EB and abamectin stimulate 510 reproduction in BPH. Thus, insecticide-enhanced reproduction is likely a key factor in 511 512 the BPH resurgence observed when farmers use EB and abamectin to control leaffolders in rice crops in China [26]. Although this is the first report of sublethal doses 513 of avermectins enhancing insect fecundity, our findings are consistent with previous 514 studies which have shown that certain insecticides, herbicides and fungicides stimulate 515 BPH reproduction [3, 7, 14, 60-65]. Intriguingly, we show that EB only induces fecundity 516 in female adults and is specific to BPH, with EB exposure failing to enhance 517 reproduction in two related species, the small brown planthopper, L. striatellus and the 518 white backed planthopper, S. furcifera, or the model insect D. melanogaster. Thus, the 519

520 mechanisms underpinning this trait appear to act exclusively on female BPH and may
521 be specific to this species.

522 Pesticides may stimulate insect reproduction through a variety of physiological and molecular mechanisms. Our data reveal that exposure to sub-lethal concentrations of 523 EB results in profound changes to female BPH fitness, leading to increases in female 524 weight, total protein content, cholesterol content, sugar content and egg production 525 and decreases in duration of the preoviposition period. Some of these findings exhibit 526 parallels with previous studies, which demonstrated that treating third-instar BPH 527 528 nymphs with either deltamethrin, triazophos, or imidacloprid led to increased soluble sugar levels in the corresponding adults [66]. Such metabolites provide the energy that 529 drives BPH reproduction and resurgence [3]. Thus, together with prior work, our results 530 531 suggest that pesticides associated with resurgence stimulate nutritional intake in BPH to fuel enhanced energy-intensive reproduction. 532

533

534 The JH signaling pathway plays an essential role in EB-induced fecundity in BPH

JH is a pleiotropic hormone which plays important roles in development and reproduction in insects [19, 67]. Circulating JH titers are regulated by factors that control JH production in the corpora allata including biosynthetic enzymes and catabolic enzymes that regulate JH levels. Our results show that EB increases circulating JH III titers in BPH females over 2–4 days after eclosion (DAE) and promotes ovary development. Previous studies have reported that triazophos and deltamethrin treatments also lead to increased circulating JH III titers in BPH females over 1–3 days post emergence. Similarly, jinggangmycin treatments were found to lead
to increased JH titers (by approximately 45–50%) in BPH females over two days post
emergence [68]. Thus our findings, in combination with these previous studies,
demonstrate that insecticide treatments can have dramatic effects on the regulation of
key insect hormones involved in pest reproduction which can in turn drive pesticide
resurgence.

Although increased JH titers following pesticide exposure have been correlated 548 with reduced levels of active JH esterase during the first three days PE [6], the type 549 550 and number of mechanisms mediating the observed increase in hormone titer has remained an open question. Our data reveal that elevated JH titer in EB-exposed BPH 551 is associated with the upregulation of genes that encode biosynthetic enzymes for JH 552 553 (JHAMT) and downstream signaling genes that can induce vg gene expression (met and *kr-h1*). Using RNAi we provide functional evidence of the role of these genes in 554 the regulation of JH III and fecundity of female BPH, and demonstrate that EB can 555 556 restore the reduction in egg production resulting from the knockdown of JHAMT, met and *kr-h1* expression. 557

JHMAT is an enzyme that catalyzes the conversion of inactive precursors of JH to active JH in the final stages of JH biosynthesis [33, 35]. Interestingly, while it has not been previously implicated in pesticide resurgence, treatment of the stored product pest *Sitotroga cerealella* with diallyl trisulfide, an insecticidal compound in garlic essential oil, was found to increase JHAMT mRNA levels [69]. Because JHMAT is the key rate-limiting enzyme in regulation of JH titer our results suggest that its enhanced 564 expression is a key molecular mechanism of pesticide resurgence in BPH.

Met is a ligand-activated member of the basic helix-loop-helix Per/Arnt/Sim 565 (bHLH-PAS) transcription factors and is the intracellular receptor for JH [36, 37]. Kr-h1 566 is a zinc finger protein that acts downstream of Met and is expressed in response to 567 JH signaling. Although the genes encoding these proteins have not been previously 568 linked to pesticide resurgence, our finding that they are upregulated following EB 569 exposure, and demonstration of their role of in promoting fecundity, is consistent with 570 previous studies. Specifically, treatment of BPH with JH III or the insecticidal analogues 571 572 methoprene or pyriproxifen was found to induce the expression of Kr-h1 [70]. Furthermore, knockdown of *Met* and *Kr-h1* in BPH brachypterous females was found 573 to result in delayed ovariole development and this was significantly more pronounced 574 575 than the response observed in BPH treated separately with dsNIMet or dsNIKr-h1 [71]. This finding provides evidence of a possible interaction between *Met* and *Kr-h1* and, 576 in combination with our data, suggests that *Met* and *Kr-h1* may act in concert to 577 578 mediate EB-enhanced fecundity.

579

580 EB-induced fecundity in BPH is dependent on the allatostatin signaling pathway

In addition to regulatory proteins that promote JH production, insects have peptides that inhibit JH biosynthesis. These include the allatostatins: FGLamides (FGLa; AstA), the W(X)6Wamides (AstB), and the PISCFs (AstC) [47, 48, 54, 55, 72, 73]. Interestingly, our results showed that EB exposure results in the marked downregulation of the expression of the genes encoding allatostatin *AstA* and its

receptor AstAR. We provide clear evidence of the functional impact of this on JH 586 synthesis and BPH fecundity by: i) demonstrating that RNAi knockdown of AstAR 587 expression results in increased JH titer and enhanced female egg production, and, ii) 588 showing that injection of female BPH with synthetic AStA peptide reduces JH titer and 589 decreases egg production. Thus, our data provide unequivocal evidence that AstA is a 590 key inhibitor of JH production in BPH. This finding is consistent with previous work 591 which has shown that FGLa/ASTs inhibit JH biosynthesis in cockroaches, and termites 592 [43, 74]. To our knowledge, our study is the first report of insecticides inhibiting the 593 594 expression of the neuropeptide receptor, AstAR, and linking this to increases in JH titer and enhanced reproduction in insects. 595

Interestingly knockdown of AstAR resulted in significant increases in the 596 597 expression of genes involved in JH synthesis/signaling including FAMeT and JHAMT *Met* and *Kr-h1*. Related to this finding, previous work has shown that knockdown of the 598 AstA receptor gene, Dar-2, in D. melanogaster results in changes in the expression of 599 600 genes encoding Drosophila insulin-like peptides (DILPs) and adipokinetic hormone 601 (AKH) signaling proteins [75]. Together with our findings, this demonstrates that AstA receptors may modulate the expression of numerous downstream genes involves in 602 metabolism, energy store and reproduction. In the case of pesticide resurgence our 603 604 results imply significant cross-talk in the expression of genes that inhibit JH production and those that promote it. 605

606

607 The GluCl plays an essential role in EB-induced fecundity in BPH

EB and abamectin are allosteric modulators of GluCls [57-59]. Our data revealed 608 that EB exposure modifies expression of the GluCl in BPH, and knockdown of GluCl 609 610 expression resulted in a reduction in both JH levels and egg production. Interestingly, the GluCl has been reported to inhibit the biosynthesis of JH in the cockroach, 611 Diploptera punctata [76]. Recent work has also reported that modulation of 612 glutamatergic signals may contribute to the photoperiodic control of reproduction in 613 bean bug, *Riptortus pedestris* [77]. Interestingly, work on *D. punctata* has revealed that 614 application of the GluCl channel agonist ivermectin, which like EB belongs to the 615 616 avermectin family, caused a decline in JH synthesis in corpus allatum glands [76]. While the inhibitory effect of ivermectin observed in this previous study differs from the 617 activating effect of EB we observed in our study, it is consistent with our finding of a 618 619 role for GluCl channel in the regulation of JH regulation. Interestingly, we found that knockdown of GluCl gene expression results in the down-regulation of JHAMT, Met 620 and Kr-h1, further revealing significant convergent relationships between genes 621 622 underpinning pesticide resurgence.

623

624 Conclusion

Our study has revealed a diverse suite of genes that act in combination to enhance JH titer and thus fecundity following BPH exposure to EB. A schematic of how these factors promote ovary development in the adult stage of *N. lugens* through the JH signaling pathway is provided in Figure 6. Our findings provide the foundation for further work to understand exactly how these genes interact and the mechanisms by

which their expression is activated or repressed by EB. Furthermore, our findings 630 provide fundamental insights into the molecular response in insects to xenobiotic 631 632 stress and illustrate that pesticides can have unexpected and negative impacts on pest populations. In this regard our findings also have applied implications for the control of 633 a highly damaging crop pest. Previous studies have reported that avermectins such as 634 abamectin are toxic to the wolf spider Pardosa pseudoannulata, which is the main 635 predator of BPH in rice crops [78, 79]. Thus, these insecticides both stimulate 636 reproduction in BPH while killing their natural enemies providing a 'perfect storm' for 637 638 damaging BPH outbreaks. Based on these findings, to avoid BPH resurgence, we suggest that the GluCls, EB and abamectin, should not be (or rarely be) applied to rice 639 640 plants at growth stages when BPHs are present. On a more optimistic note, our 641 findings have identified numerous genes that play key roles in BPH reproduction and thus represent promising targets for the development of novel controls against this 642 important pest. 643

645 Figure 6



646

647 Figure 6. Schematic of the proposed regulatory pathway of EB-enhanced fecundity in BPH. Emamectin benzoate (EB) exposure results in the upregulation of genes that promote 648 649 JH production (JHAMT, Met and Kr-h1) and the downregulation of genes that inhibit it 650 (allatostatin, AstA and allatostatin A receptor, AstAR). This transcriptome reprograming is 651 dependent on the action of EB on its molecular target the glutamate-gated chloride channel 652 (GluCI) receptor. The resulting increased JH titer promotes vg synthesis and increased fecundity in EB exposed insects. We observe significant cross-talk in the expression of genes 653 that inhibit JH production and those that promote it, with AstAR inhibiting the expression of 654 JHAMT, Met and Kr-h1 and GluCl activating the expression of JHAMT which is responsible for 655 JH synthesis, and the JH signalling downstream genes Met and Kr-h1. 656

657 Materials and methods

658 Insects

BPH was initially collected from Wanan, JiangXi Province in 2020, reared on 'Taichung Native 1' (TN1) rice seedlings in the laboratory without exposure to any insecticides. The strain was maintained in a climatic chamber at $27 \pm 1^{\circ}$ C, with relative humidity of 70 ± 10% and a light: dark = 16 h: 8 h photoperiod.

663 Chemicals

Emamectin benzoate (95.2%) was obtained from Hebei Weiyuan Co., Ltd. (Hebei,
China). Abamectin (96.8%) was obtained from Hebei Weiyuan Co., Ltd. (Hebei, China).
Pyriproxyfen (98%) was obtained from Shanghai Shengnong Co., Ltd. (Shanghai,
China). Methoprene (S)-(+) (mx7457-100mg) was purchased from Shanghai MaoKang
Biotechnology Co., Ltd., (Shanghai, China). Juvenile hormone standard sample
(J912305-10mg) was purchased from Shanghai Macklin Biotechnology Co., Ltd.,
(Shanghai, China).

671 Bioassay

Different life stages of insects were used to perform bioassay to investigate the effects of insecticide on nymphs and adults. To test whether treatment of the nymph stage of insects would promote reproduction in female, we used 4th instar nymphs of BPH or 3rd instar nymph of *Laodelphax striatellus* and *Sogatella furcifera* to perform bioassays. To test whether treatment of the adult stage of insects would promote reproduction in female, we used newly emerged male and female BPH.

678 **Systemic route:** The rice-seeding dipping bioassay method was used to evaluate

the susceptibility of BPH, L. striatellus and S. furcifera to EB. Technical-grade 679 insecticides were dissolved in acetone as stock solution then diluted in a series of six 680 681 concentrations with water containing 0.1% Triton. Selected rice seedlings at the 6-8 cm growth stage were dipped in insecticide solutions for 30 s and then air-dried at 682 room temperature. The roots of the rice seedlings were wrapped with cotton strips and 683 placed seedlings placed in a plastic cup 5 cm in diameter. Fifteen insects were 684 introduced into each plastic cup for each replicate. The top of the cup was then selaed 685 with gauze to prevent escape. All experiments comprised at least three biological 686 replicates. Control rice seedings were treated with 0.1% Triton X-100 water solution 687 only. All treatments were maintained under standard conditions of 27 ± 1 °C and 70-688 80% relative humidity with a 16 h light/8 h dark photoperiod. Mortality was assessed 689 690 after 4 d for N. lugens or 2 d for L. striatellus and S. furcifera after treatment with insecticides. The insects were considered dead if they were unable to move after a 691 gentle prodding with a fine brush. 692

693 For Drosophila larvae bioassay, we adopted a method described previously in our 694 lab with minor modifications [80]. Briefly, twenty third instar larvae were placed in fly vials containing fly food (based on corn powder, brown sugar, yeast and agar) 695 supplemented with EB of different concentrations. Four concentrations (LC₁₀, LC₃₀ and 696 697 LC₅₀) were tested together with a negative (no insecticide) control. For Drosophila adult bioassays, we selected virgin females three days after eclosion. Several 698 699 concentrations were overlaid onto fly food in standard Drosophila vials and allowed to dry overnight at room temperature. 15 adult flies (three days after eclosion) were then 700

added to each vial and mortality assessed after 2 d. Four replicates were carried out
for each concentration. Control mortality was assessed using vials containing food with
solvent minus insecticide.

Contact route: For topical bioassays working insecticide solutions were prepared 704 705 in acetone. 4th instar nymphs or newly emerged males/females were anesthetized with carbon dioxide for 5 s, and then 0.04 µl/insect test solution applied topically to the 706 dorsal plates with a capillary micro-applicator [26]. Insects were then placed in an 707 708 artificial climate incubator with a temperature of 27±1°C, a photoperiod of 16:8 h (L:D), 709 and a humidity of 70%±10%. Mortality was determined 2 d after treatment. Data with over 20% mortality in the control treatment were discarded, and the assay was 710 repeated at least three times. 711

712 **Fecundity assays**

Fourth instar nymphs of BPH and 3rd instar nymph of *L. striatellus* and *S. furcifera* 713 were treated with the LC_{15} and LC_{50} of EB, and then transferred to fresh rice seedlings. 714 After eclosion, the adults were used in the following experiment. Newly emerged 715 treated adults and untreated adults were paired to produce four groups: untreated 716 males and untreated females (*d*ck×Ωck; ck indicates untreated); untreated males and 717 treated females (3 ck× 2t; t indicates insecticide treated); treated males and untreated 718 females ($\partial t \times \Omega ck$); treated males and treated females ($\partial t \times \Omega t$). Each group comprised 719 720 at least 10 mating pairs. All pairs were transferred to glass cylinders (diameter 5 cm 721 and height 20 cm) containing rice plants (25 days old almost 20 cm high) as a food source for eleven days. The number of eggs and nymphs in plants were counted by a 722

723 microscope.

For *Drosophila* egg-laying assay, we adopted our previous method [81]. Briefly, insecticide-treated virgin females were paired with untreated males for three days and then the mated females transferred into the *Drosophila* ovipositing apparatus. The number of eggs were counted after 16 hours.

728 Fitness analysis

The fitness of EB-treated BPH were analyzed using methods reported previously 729 [10]. We selected two groups, ($\partial ck \times Q ck$) and ($\partial ck \times Q t$), to study the effects of the LC₅₀ 730 concentration of EB on BPH fitness. In the case of systemic exposure, 4th instar 731 nymphs of BPH were treated with the LC₅₀ of EB for 4 days and then transferred to 732 tubes containing untreated rice plants for individual rearing. The rice plants were 733 734 replaced every three days with untreated plants. The emergence ratio, female ratio, preoviposition period, female longevity, brachypterism female ratio and female weight 735 were calculated. 736

737 Examination of ovary development

Adult females from $\partial ck \times Q ck$ control and $\partial ck \times Q t$ group on 1, 3, 5, 7 DAE were dissected for observe the ovary development. The detained eggs in ovary were photographed and recorded. Each group has at least fifteen replicates.

To examination whether EB treated impairs egg maturation, we dissected untreated or EB-treated ovaries and fixed them in 4% paraformaldehyde (PFA) in phosphate buffered saline (PBS) for 30 min at room temperature. After four washes of

744	10 min (4 × 10 min) in PAT3 (PBS with 0.5% Triton X-100 and 0.5% bovine serum
745	albumin), the ovaries were then incubated with DAPI (4',6-diamidino-2-phenylindole,
746	100 nM) and Actin-stain 670 Fluorescent Phalloidin (200 nM). Imaging was performed
747	using Zeiss LSM980 confocal laser microscope.
748	Measurements of glycogen, triglyceride, total protein content, cholesterol and
749	four sugars
750	The content of glycogen, triglyceride, cholesterol and total protein was
751	determined by spectrophotometry at 620 nm, 510 nm, 510 nm and 562 nm
752	respectively using the glycogen assay kit (A043-1-1), triglyceride reagent kit (A110-1-
753	1), cholesterol assay kit (A111-1-1) and total protein assay kit (A045-2-2) obtained
754	from Nanjing Jiancheng Bioengineering Institute following to the manufacturer's
755	instructions. The determined results were normalized to the protein content in the
756	sample, which was determined using BCA Protein Assay Reagent Kit (Thermo
757	Scientific, Waltham, USA). Each sample contained tissue extracts from five adult
758	female BPH, with three biological replicates per sample.
759	To assess the content of four sugars (sucrose, glucose, fructose and trehalose)
760	in the extract of BPH tissue, the same extraction method was used as above. Sugar
761	content was quantified using the colorimetric method by scanning at 290 nm, 505
762	nm, 285 nm and 620 nm respectively using the sucrose reagent kit (A099-1-1),
763	glucose reagent kit (F006-1-1), fructose reagent kit (A085-1-1) and trehalose reagent
764	kit (A150-1-1) obtained from Nanjing Jiancheng Bioengineering Institute based on the
765	manufacturer's instructions. Each sample contained tissue extracts from five adult

female *N. lugens*, with three biological replicates per sample.

767	Determination of Juvenile hormone III and ecdysone titers of BPH by ELISA
768	The titer of Juvenile hormone III in BPH was measured using the Juvenile
769	hormone ELISA Kit (Lot Number: HLE92086, Shanghai HaLing Biotechnology Co., Ltd.,
770	Shanghai, China) which employs the dual-antibody sandwich ELISA method. The titer
771	of ecdysone in BPH were measured using the ecdysone ELISA Kit (Lot Number:
772	ZK14705, Shanghai ZhenKe Biotechnology Co., Ltd., Shanghai, China). At least three
773	biological replicates were employed for each treatment.

774 Determination of Juvenile hormone III titer in BPH using HPLC-MS/MS

The whole bodies of 5 individuals BPH were mixed with 1 ml of n-hexane, followed 775 776 by centrifugation at 10,000×g for 10 min, the upper hexane layer was then dried with nitrogen, dissolved in methanol and sonicated for 10 min, after centrifugation at 777 10000×g for 10 min, the supernatant was collected through the organic filter membrane 778 of 0.22 µm into 2 ml vials for JH III determination. JH III standard sample (J912305-779 10 mg) purchased from (Shanghai McLean Biochemical Technology Co. Ltd), 780 dissolved in methyl alcohol as stock solution 10,000 mg/L was diluted in a series of six 781 782 concentration gradients to serve as a reference. Liquid chromatography-tandem mass spectrometry (LC-MS/MS) was then carried out using UPLC Xevo TQ-S Micro (Waters 783 technology), quantitative method according to the external standard, the 784 785 chromatographic column was EC-C18 (4.6 mm×150 mm, 2.7 µm), column temperature was 30°C, injection volume was 20 µl, elution flow rate was 0.3 ml/min, and the mobile 786

- 787 phase was acetonitrile:formic acid water (90:10), detection wavelength was 218 nm,
- the peak height was used for quantification.

789 Cloning, sequence and phylogenetic analysis

The NCBI database and BLAST program were used to carry out sequence 790 alignment and analysis. Open Reading Frames (ORFs) were predicted with EditSeq. 791 Primers were designed using the primer design tool in NCBI. Total RNA Extraction was 792 793 extracted from 30 adults BPH using TRIzol reagent (Invitrogen, Carlsbad, CA, USA) 794 according to the manufacturer's instructions. cDNA was synthesized using the Biotech M-MLV reverse transcription kit. Full-length gene sequences were amplified by PCR 795 using cDNA as template and Rapid Tag Master Mix (Vazyme Biotech, Cat# P222-02). 796 797 The PCR product was purified on a 1% agarose gel, cloned into pClone007 Simple Vector Kit (Tsingke Biotech, Cat# TSV-007S), and then sequenced using the 3730 XL 798 DNA analyzer (Applied Biosystems, Carlsbad, CA, USA). Table S2 contains a list of 799 800 the primers used in this study.

The exon and intron architectures of AT, AstA, AstB, AstCC and AstACCC were 801 predicted based on the alignments of putative ORFs against their corresponding 802 genomic sequences. Sequence similarity/annotations and orthologous gene searches 803 804 were performed using BLAST programs available in NCBI. Multiple alignments of the complete performed amino acid sequences were with Clustal Omega 805 806 (http://www.ebi.ac.uk/Tools/msa/clustalo). Phylogeny was conducted using the maximum likelihood technique to create phylogenetic trees and these were 807

808 bootstrapped with 1000 replications were used using the MEGA 6 software [82].

809 RNA interference

810	Double-stranded RNA (dsRNA) of <i>gfp</i> (green fluorescent protein), <i>JHAMT</i>
811	(juvenile hormone acid O-methyltransferase,), Met (methoprene-tolerant), Kr-h1
812	(krüppel homolog 1), AstAR (allatostatin-A receptor) and GluCl (glutamate-gated
813	chloride channel) was prepared using Ambion's MEGAscript T7 kit instructions
814	following the manufacturer's instructions. The primer sequences for double-stranded
815	RNA synthesis are listed in Table S2. Newly emerged females were injected with 40
816	nl (5,000 ng/µl) of double-stranded RNA of <i>gfp</i> (<i>dsgfp</i>) or double-stranded RNA of the
817	target genes in the conjunctive part between prothorax and mesothorax of insects. In
818	the RNAi experiments, BPH were then treated with the LC_{50} of EB 24h after dsRNA
819	injection and the whole body sampled for qRT-PCR analysis.

820 Quantitative RT-PCR

821 Fourth instar nymphs of BPH were treated with EB after which total RNA was extracted from 5th instar nymphs and 1-7 day post-eclosion females of *N. lugens* using 822 the methods detailed above. The HiScript® II RT SuperMix for qPCR (+gDNA wiper) 823 kit from Vazyme, Nanjing, China, was used to create first-strand cDNA. Primer3 824 (http://bioinfo.ut.ee/primer3/) was used to design real-time quantitative PCR (qPCR) 825 primers listed in Table S2. mRNA levels of candidate genes were detected by gPCR 826 827 using the UltraSYBR Mixture (with ROX) Kit (CWBIO, Beijing, China). Each reaction contained 2 µL of cDNA template (500 ng), 1 µL each forward and reverse qPCR primer 828

(10 µM), 10 µL of UltraSYBR mixture buffer, and 6 µL of RNase-free water. Q-PCR 829 was run on an ABI 7500 Real-Time PCR System (Applied Biosystems, Foster City, CA) 830 831 under the following conditions: 5 min at 95°C, followed by 40 cycles of 95°C for 10 s and 60°C for 30 s. Three independent biological replicates and four technical replicates 832 were used in each qPCR experiment. The housekeeping genes 18S ribosomal RNA 833 of BPH were selected to normalize the expression of candidate genes. The $2^{-\Delta\Delta Ct}$ 834 method (Ct represents the cycle threshold) was used to measure relative expression 835 levels [83]. Three biological replicates were used for statistical comparison between 836 837 samples. Table S2 contains a list of the primers used in this study.

838 Statistics

PoloPlus v2.0 (LeOra Sofware 2008) was used to calculate the lethal 839 concentration (LC₅₀) and 95% fiducial limits (95% F.L.). GraphPad Prism 8.0 software 840 (GraphPad Software Inc., San Diego, USA) was used to generate graphs and perform 841 statistical analysis of data. Data presented in this study were first verified for normal 842 distribution using the D'Agostino-Pearson normality test. One-way analysis of 843 variance (ANOVA) with Duncan's multiple range test was used to test differences 844 among multiple groups of normally distributed groups of data. Student's t test was used 845 to test the differences between two groups. If not normally distributed, Mann–Whitney 846 test was used for pairwise comparisons, and Kruskal-Wallis test was used for 847 comparisons among multiple groups, followed by Dunn's multiple comparisons. All 848 849 data are presented as mean ± s.e.m. The sample sizes and statistical tests used for each experiment are stated in the figures or figure legends. 850

851

Table S2 Sequences of oligonucleotide primers used in this study.

Primers	Primer sequences (5' - 3')
For cDNA cloning	
Allatotropin:	
NIAT-F	TACGCGGCCAAACACACTTA
NIAT-R	AGGGAAAGAGGGCGAAATTCA
Allatostatins:	
NIAstA-F	TCGGCCGTCACAAGTCAAG
NIAstA-R	CCGAACCCGTACTTCATGCT
NIAstB-F	ACCGGGCTCACAGGAATTTT
NIAstB-R	TGTAGGCGCAGATCTTGAGG
NIAstCC-F	AACACAGCTCTACGAGGCAC
NIAstCC-R	CCAAGCAGGTGACTGCCATA
NIAstCCC-F	TTTGTGTGTGCTTGCAGGTG
NIAstCCC-R	GGATAGAAACGGTAGATTTGGTAGA
Allatostatins receptor:	
NIA16-F	CCTCATTGTGGAACCACCGA
NIA16-R	CGCAGCTGTAAGGTGGAAGA
NIA2-F	GAACGTAATGGGAGTCGGCA
NIA2-R	GTTTTTGTGAGCGCCGACTT
NIA10-F	ATGCAAAACACGGCCAGCCT
NIA10-R	TTAATCGTCTCTGCTCAACTCCAAAGGAAGGT
NIA1-F	CGACCAGACCACTCTACTGC
NIA1-R	ACGTGGACCTCACTATACCAAAAA
For Quantitative RT-PCR	
Q-NI18S-F	CGCTACTACCGATTGAA
Q-NI18S-R	GGAAACCTTGTTACGACTT
Q-vitellogenin-F	GTGGCTCGTTCAAGGTTATGG
Q-vitellogenin-R	GCAATCTCTGGGTGCTGTTG
Q-Vitellogenin receptor-F	AGGCAGCCACAGATAACCGC
Q-Vitellogenin receptor-R	AGCCGCTCGCTCCAGAACATT
Q-JHE-F	GAGCCTCACATCCACAGC
Q-JHE-R	AATGGGAGCCCTACGC
Q-NIMet-F	GGTGGTAAACGGATTGGAAA
Q-NIMet-R	CATCGTCAGCCAACTCGATA
Q-NIFAMet-F	GCAAAGTCAGCAATCCGCAAGAAC
Q-NIFAMet-R	ACACCGTAGTGGGTGACAACGAATG
Q-JHAMT-F	GAACCTGCAGGCCAAACACA
Q-JHAMT-R	ACCACTCGGTTGGGCTGAAT
Q-NIKr-h1-F	TGATGAGGCACACGATGACT
Q-NIKr-h1-R	ATGGAAGGCCACATCAAGAG
Q-NIAT-F	CACGATACGTGGCTTCAAGA
Q-NIAT-R	ACGATCACTTTCGCCAATTC
Q-NIAstA-F	AGGACTTACTGGGCGAGGAT

Q-NIAstA-R	GGTGTCTCGTTTCCTGGTGT
Q-NIAstB-F	AGCGAGCTAGACGAGGACAA
Q-NIAstB-R	TCGTCTCTGCTCAACTCCAA
Q-NIAstCC-F	CTGCTCCCAGTGAAAAGGAG
Q-NIAstCC-R	GCTTCCAGTAACTGCGCTTC
Q-NIAstCCC-F	TGTGCTTGCAGGTGGTAGTC
Q-NIAstCCC-F	AGAAGCATGTGACTGCGTTG
Q-NIA2-F	TCCTGGTGCTGAAGAGTGTG
Q-NIA2-R	CTTTTCGGGCCCATTAATTT
Q-NLA1-F	ATATCGGCACCGAAGATGAG
Q-NLA1-R	GTCTGACCCGACAGGTTCTC
Q-NLA10-F	ACTGGGTGTCGACCAATCTC
Q-NLA10-R	TCGGTAGCGAGGAAGACAGT
Q-NLA16-F	TACCGTTCTGTGGGATGTCA
Q-NLA16-R	CCGGGATATCAAAGACGAGA
Q-NIGluCI-F	CACTGACTGAGGCCAACAGA
Q-NIGluCI-R	GCTGGCCATTCTTAGTGAGC

For double-stranded RNA synthesis

T7-EGFP-F	TAATACGACTCACTATAGGGCGTAAACGGCCACAAGTTCA
T7-EGFP-R	TAATACGACTCACTATAGGGGACTGGGTGCTCAGGTAGTG
T7-Kr-h1-F	TAATACGACTCACTATAGGGCGCCAGTGAAAGTGAGACCT
T7-Kr-h1-R	TAATACGACTCACTATAGGGGAGACCGCAAGTGGTTCTGA
T7-Met-F	TAATACGACTCACTATAGGGCCACCAACCAGCAGATGAACCTGA
T7-Met-R	TAATACGACTCACTATAGGGCCACGCAAAGCCTCGTACTCTTGG
T7-JHAMT-F	TAATACGACTCACTATAGGGCTCCAGGCCATTGTCCCTCA
T7-JHAMT-R	TAATACGACTCACTATAGGGTTGGCCTGCAGGTTCTTTGG
T7-AstA-R-F	TAATACGACTCACTATAGGGTACTGCCGTTCTGGCCTTTT
T7-AstA-R-R	TAATACGACTCACTATAGGGGTGATCTGGAAGAGCGGCTT
T7-Glucl-F	TAATACGACTCACTATAGGGACACATCACCTGCTCACCTG
T7-Glucl-R	TAATACGACTCACTATAGGGGTGTGTTTGCCTGCTGTCTG

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