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Pan, D, Luo, Q-J, O Reilly, AO, Yuan, G-R, Wang, J-J and Dou, W (2023) Mutations of voltage-gated sodium channel contribute to pyrethroid resistance in *Panonychus citri*. *Insect science*. pp. 1-14. ISSN 1672-9609

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1 *Manuscript submitted to Insect Science*

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3 **Mutations of voltage gated sodium channel contribute to pyrethroid**
4 **resistance in *Panonychus citri***

5

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28

29 **Abstract**

30 Insecticide resistance in *Panonychus citri* is a major obstacle to mite control in
31 citrus orchards. Pyrethroid insecticides are continually used to control mites in China,
32 although resistance to pyrethroids has evolved in some populations. Here, the
33 resistance to the pyrethroid fenpropathrin was investigated and seven out of eight
34 field-collected populations of *P. citri* exhibited high level of resistance, ranging from
35 171- to 15,391-fold higher than the susceptible (SS) comparison strain. Three voltage-
36 gated sodium channel (VGSC) mutations were identified in the tested populations:
37 L1031V, F1747L, and F1751I. Amplicon sequencing was used to evaluate the
38 frequency of these mutations in the nineteen field populations. L1031V and F1751I
39 were present in all populations at frequencies of 11.6–82.1% and 0.5–31.8%,
40 respectively, while the F1747L mutation was only present in twelve populations from
41 Chongqing, Sichuan, Guangxi and Yunnan provinces. Introduction of these mutations
42 singly or in combination into transgenic flies significantly increased their resistance to
43 fenpropathrin and these flies also exhibited reduced mortality after exposure to the
44 pyrethroids permethrin and beta-cypermethrin. *P. citri* VGSC homology modeling and
45 ligand docking indicate that F1747 and F1751 form direct binding contacts with
46 pyrethroids, which are lost with mutation, whereas L1031 mutation may diminish
47 pyrethroid effects through an allosteric mechanism. Overall, the results provide
48 molecular markers for monitoring pest resistance to pyrethroids and offer new
49 insights into the basis of pyrethroid actions on sodium channels.

50 **Key words:** *Panonychus citri*, kdr mutation, pyrethroid resistance, amplicon
51 sequencing, transgenic flies, molecular docking

52

53 **Introduction**

54 Voltage-gated sodium channels (VGSCs) are critical components of nearly all
55 animal nervous systems (Banazeer et al., 2022). VGSCs conduct sodium ions across
56 the plasma membrane to initiate and propagate electrical signals that regulate insect
57 behavior (Williams et al., 2022). Pyrethroids target the VGSCs of insects by
58 stabilising the open functional state, resulting in nervous system dysfunction. They
59 are highly efficient and exhibit specific selective toxicity for insects over mammals,
60 leading to their widespread use for terminating or controlling pests or mites (Scott,
61 2019). However, the extensive use of pyrethroid insecticides has promoted the
62 evolution of resistance, thereby reducing the efficacy of currently used pyrethroids
63 (Pan et al., 2020). The presence and evolution of resistance to pyrethroids has been
64 previously investigated in *Musca domestica* (Scott, 2017, Roca-Acevedo et al., 2023),
65 *Rhipicephalus microplus* (Kumar et al., 2020), *Leptinotarsa decemlineata* (Molnar &
66 Rakosy-Tican, 2021), *Plutella xylostella* (Banazeer et al., 2022), and mosquito vectors
67 (Scott et al., 2015, Smith et al., 2016, Amelia-Yap et al., 2018, Chen et al., 2020).

68 Pyrethroids stimulate nerve cells to prolonged-activation and consequently leads
69 to paralysis or death, which is known as *knockdown* (Davies et al., 2008). An
70 important resistance mechanism against pyrethroids or DDT is known as ‘knockdown
71 resistance’ (*kdr*), involving various mutations found in the VGSC of many pests or
72 mites (Rinkevich et al., 2013). More than 61 sodium channel mutations, or
73 combinations of mutations, have been reported to be responsible for, or associated
74 with, pyrethroid resistance in various arthropods (Dong et al., 2014). Recently, several
75 *kdr* mutations conferring pyrethroid resistance have been identified in pest mites. For
76 example, four amino acid substitutions in VGSC (F1528L, L1596P, I1752V, and
77 M1823I) have been determined to be associated with fluvalinate resistance in the
78 varroa mite (*Varroa destructor*) (Wang et al., 2002). Further, the M918T mutation of
79 the domain II region led to high resistance to bifenthrin in *Tetranychus evansi* (Nyoni
80 et al., 2011). In addition, the mutations L1024V, A1215D, and F1538I have been
81 detected in a pyrethroid-resistant strain of *Tetranychus urticae*, with the L1024V and
82 A1215D mutations potentially contributing to fenpropathrin resistance (Kwon et al.,

83 2010). Moreover, the point mutation F1538I at the IIS6 site of the sodium channel
84 has also been identified in a fenpropathrin-resistant population of *Panonychus citri*
85 (Ding et al., 2015).

86 *Panonychus citri* (McGregor) (Acari: Tetranychidae), also known as the citrus red
87 mite, is a dominant pest mite in global citrus orchards. The mites feed on the leaves,
88 fruits, and the fresh shoots of various citrus plants, causing severe damage to the
89 plants (Alavijeh et al., 2020). Defoliation and botchy fruits appear upon profusive
90 mite outbreaks in orchards. Fenpropathrin is a pyrethroid chemical that has been
91 extensively used for mite control in many Chinese citrus orchards (Hu et al., 2010).
92 However, *P. citri* has already developed high resistance to fenpropathrin in several
93 citrus production areas (Pan et al., 2020). Nevertheless, the frequency of VGSC
94 mutations in *P. citri* across these large areas and the contributions of single mutations,
95 or combinations, to pyrethroid resistance have not yet been explored.

96 Here, resistance to fenpropathrin was investigated among several *P. citri*
97 populations collected from five provinces in China. Three amino acid substitutions of
98 VGSC were identified in fenpropathrin-resistant populations and the frequencies of
99 these mutations in nineteen *P. citri* field populations were subsequently analyzed. The
100 roles of the VGSC mutations in *P. citri* pyrethroid resistance were further analyzed by
101 constructing transgenic flies carrying these mutations as single mutations or
102 combinations. Further, a homology model of *P. citri* VGSC was generated to
103 investigate differences between wild type (WT) and mutated models that bind to
104 pyrethroids.

105 **Materials and methods**

106 **Mites**

107 A relatively susceptible strain (SS) of *P. citri* was collected from citrus orchards at
108 the Citrus Research Institute at Southwest University in Chongqing, China (Pan et al.,
109 2020). Mites were maintained at $27 \pm 1^\circ\text{C}$, $60 \pm 5\%$ relative humidity, and with a 16:
110 8 h light / dark photoperiodic cycle. The SS strain was used as the control for toxicity
111 bioassays. Eight field populations of *P. citri* were collected for bioassays from the
112 Yunnan, Jiangxi, Chongqing, Sichuan, and Guangxi provinces. Sampling location

113 details are provided in Table S1. Adult female mites were collected for each
114 population from at least 10 trees per citrus orchard. Over 5,000 adult female mites
115 were collected per field population and randomly selected for bioassays. In addition,
116 nineteen field populations were collected for DNA extraction and sequencing to
117 investigate mutation frequency (Table S2). All mites collected from orchards were
118 sampled as described above. Of the collected mites, 100 adult female mites were
119 preserved in ten 1.5 mL centrifuge tubes and then preserved in 100% ethanol at room
120 temperature.

121 **Bioassay chemicals**

122 The acaricide chemical 20 g/L EC fenpropathrin (Well-Done Chemical,
123 Hangzhou, China) was used for bioassay. Other chemicals were used to test
124 insecticide susceptibility in transgenic *D. melanogaster*, including permethrin (type I,
125 95% cis-trans isomer mixture: Macklin Inc, Shanghai, China), beta-cypermethrin
126 (type II, 95%; Xiya Reagent, Shangdong, China), and fenpropathrin (92.9%; Veyong
127 Animal Pharmaceutical Co., Ltd, Hebei, China).

128 **Bioassays**

129 The susceptibility to fenpropathrin in field populations of *P. citri* were assayed
130 using the leaf-dipping method, as described by Yamamoto et al (Yamamoto et al.,
131 1995). Leaf discs of 25 mm diameter were produced from thoroughly washed fresh
132 sweet orange (*Citrus sinensis* (L.) Osbeck) leaves and placed on water-saturated
133 sponges in Petri dishes of 9 cm diameter. A total of 25–30 females were subsequently
134 transferred to each disc with a soft brush. After 2–3 h, the leaf discs with mites were
135 dipped in acaricide solutions containing seven different concentrations for 5 s.
136 Superfluous liquid on the discs was absorbed with absorbent paper to avoid drowning
137 the mites. Leaf discs treated with distilled water were used as control. Sponges were
138 kept wet with enough water to ensure they did not dry. Each evaluated concentration
139 included five replicate assays. All leaf discs were incubated at $25 \pm 0.5^{\circ}\text{C}$, with 60%
140 relative humidity and a 16: 8 h (L:D) photoperiod. Mite mortality was assessed after
141 24 h. Drowned mites on the edge of the leaf discs and dead mites due to human
142 factors were not recorded as mortalities, while mites that could not move after light

143 touching with a camel hairbrush were scored as dead.

144 **Detection of candidate target site mutations**

145 Genomic DNA was extracted from approximately 200 adult female mites from
146 different field populations and subjected to PCR amplification. The qualities and
147 concentrations of DNA were assessed with a spectrophotometer (Nanodrop 2000,
148 Thermo Fisher, USA). PCR amplification of *P. citri* VGSC (GenBank: KF646792)
149 regions was conducted, as described for *P. citri* (Ding et al., 2015) and *Tetranychus*
150 *urticae* (Inak et al., 2019, Kwon et al., 2014), using primers shown in Table S3. PCR
151 products were purified using the MiniBEST DNA Fragment Purification Kit (TaKaRa,
152 Dalian, China) and sequenced at the Beijing Genomics Institute (BGI, Beijing,
153 China). Nucleotide or protein sequence alignments were visualized in MEGA (version
154 6.0, USA) and Jalview (version 2.0, USA).

155 **Amplicon sequencing of pooled female adults to detect mutations**

156 *DNA preparation*

157 For each group of *P. citri* adult females from citrus orchards, DNA was extracted
158 from a pool containing 80 adult female mites from each field population and more
159 than eight replicates were prepared. DNA was extracted using a Mollusc DNA Kit
160 (Omega Bio-Tek, USA), with DNA quality and quantity assessed with a
161 spectrophotometer (Nanodrop 2000, Thermo Fisher, USA).

162 *PCR amplification with tagged primers*

163 Six base-pair oligonucleotide barcodes were added to the 5' ends of the forward
164 and reverse primers (Table S4) to enable multiplex sequencing of amplicon pools and
165 subsequent identification of different field samples. PCR mixtures and amplifications
166 were used as described above. PCR products were separated by 1% agarose gel
167 electrophoresis and target DNA fragments were purified using a TaKaRa MiniBEST
168 Agarose Gel DNA Extraction Kit (TaKaRa, Dalian, China).

169 *Next-generation sequencing (NGS) of pooled amplicons*

170 PCR products from different populations from each year were pooled for NGS.
171 Amplicon concentrations per population were quantified and PCR products were
172 pooled in equimolar concentrations. Each pool corresponded to a VGSC gene

173 fragment amplicon group. Sequencing libraries were constructed using TruSeq DNA
174 PCR-Free Kits (Illumina, CA). Pooled amplicons were then sequenced on the
175 Illumina HiSeq2500 platform used paired-end 250 bp sequencing at BIOZERON
176 Biotechnology Ltd. (Shanghai, China).

177 *DNA read sorting by tags*

178 Raw sequence quality data was assessed using the FastQC software program
179 (<http://www.bioinformatics.babraham.ac.uk/projects/fastqc>). Preprocessing of raw
180 data was conducted with the FASTX_Clipper program, followed by quality filtering
181 with the FASTQ_Quality_Filter. The raw data was subsequently de-duplicated using
182 the FASTX_Collapser program (FASTX-Toolkit,
183 http://hannonlab.cshl.edu/fastx_toolkit/index.html). Clean data were mapped to the
184 reference genome of *P. citri* (accession number: GWHBAOM00000000) with the
185 BWA (Li & Durbin, 2009) aligner and then filtered using the Seqtk program
186 (<https://github.com/lh3/seqtk>). Haplotypes were then classified using the
187 FASTX_Barcode_Splitter program based on the unique barcode tags of each
188 population. A threshold value of $1/(2n)$ was used to filter the data and remove false
189 positive haplotypes from each population, where n is the number of adult females in
190 the pool and represents the theoretical minimal allele frequency for the site (Edwards
191 et al., 2018). Haplotypes were discarded when their frequency was less than the
192 threshold.

193 **Correlation of resistance level versus allele frequency in field populations of *P.***

194 *citri*

195 According to the determined resistance levels in field populations and allele
196 mutation frequencies, the correlation between the frequencies of each mutation (and
197 their combinations) and the resistance level in field populations were analyzed. The
198 resistance ratios of field populations (Table 1) were plotted against the frequency of
199 these mutations (Table 2), and linear regression lines were generated (GraphPad Prism
200 8.0 software program, GraphPad Software Inc.). When the allele frequencies at each
201 mutation site were conducted in various combinations for correlation analysis, mean
202 parameter values were used (Kwon et al., 2010, Kwon et al., 2004, Kwon et al., 2012,

203 Chang et al., 2012).

204 **Transgenic *D. melanogaster* and mortality bioassays**

205 *Transgenic expression of VGSC (WT and mutated) in Drosophila strains*

206 To confirm whether mutations of VGSC in *P. citri* conferred pyrethroid
207 resistance, the VGSC genes (6,729 bp) of WT and mutant (L1031V, F1747L and
208 F1751I) genes were expressed in *D. melanogaster* via the Gal4/UAS system from
209 Fungene Biotech (<http://www.fungene.tech/>) (Beijing, China). WT and mutant genes
210 were synthesized and the ORFs of VGSC genes were subcloned into the expression
211 vector (pJFRC28-10XUAS-IVS-GFP-p10 vector). Clones were microinjected into
212 the germ line of *D. melanogaster* (W^{1118}) embryos carrying the attP40 (25C6 of
213 chr.2) docking site on chromosome 2 to generate transgenic lines using the PhiC31
214 system. The transgenic lines were first crossed with *Bc/CyO* and screened for
215 candidates that were then verified by PCR (Table S5) and sequenced. Successful
216 construction of UAS-VGSC lines (UAS-WT, UAS-L1031V, UAS -F1747L, UAS -
217 F1751I, UAS -L1031V+ F1747L, UAS -L1031V+ F1751I, UAS-F1747L+F1751I,
218 UAS-L1031V+F1747L+F1751I) were balanced by Actin5C-GAL4/*CyO* to drive
219 ubiquitous expression. Gal4/UAS-VGSC lines were finally obtained and PCR was
220 then used to validate VGSC mutation sites (Table S3 and Fig. S1). Flies with the
221 same background as the UAS transgenic lines, but devoid of the driver Actin5C-Gal4
222 line to overexpress VGSC were used as controls. All fly lines were reared on an
223 artificial diet at 25°C, 65% relative humidity and a 12: 12 h light/dark photoperiod.

224 *Drosophila contact bioassays*

225 Bioassays were conducted to assess the susceptibility of transgenic flies to
226 fenpropathrin, as previously described (Ibrahim et al., 2015, Riveron et al., 2013).
227 Insecticide filter papers with five to seven concentrations of fenpropathrin were
228 prepared in distilled water that contained 1% acetone and 0.1% Triton-X 100. The
229 papers were rolled and placed in 5 cc glass vials. Vials were then plugged with cotton
230 soaked in 20% (wt/vol) sucrose, followed by transfer of 25 flies into each vial. The
231 mortality plus knockdown, when their legs cannot move or body back down, was
232 scored after exposure to insecticides for 24 h. Six replicates were used for each

233 concentration. All tested flies were reared at 25°C, 65% relative humidity, and with a
234 12: 12 h light/dark photoperiod. Student's *t*-tests were used to compare differences in
235 mortality and knockdown of treatment (Gal4/UAS lines) groups with each control
236 (UAS line) group, in addition to differences between mutant and WT groups.

237 In addition to the above experiments, the ability of the VGSC mutations to
238 confer resistance to two types of pyrethroids, type I (permethrin) and type II (beta-
239 cypermethrin) was evaluated. Pyrethroid filter papers (permethrin at 50 mg/L and
240 beta-cypermethrin at 5 mg/L) were prepared as described above. The mortality plus
241 knockdown was scored after exposure to insecticides for 1, 2, 3, 6, 12, 24, 36 and 48
242 h. Six replicates were used for each assay. Student's *t*-tests were used to compare
243 differences in knockdown rate between the experimental (Gal/UAS lines) and control
244 (UAS line) groups, in addition to between mutant and WT groups.

245 ***P. citri* VGSC homology modelling and ligand docking**

246 A homology model of *P. citri* VGSC was generated using the VGSC structure of
247 *Periplaneta americana* (PDB entry 5X0M) as template, as described.(Clarkson et al.,
248 2021) Sequences were aligned using Clustal Omega (Sievers et al., 2011) and the
249 MODELLER program was used to generate 50 homology models (Webb & Sali,
250 2016). The internal scoring function of MODELLER was used to select 10 models
251 that were then visually inspected and submitted to the VADAR webserver (Willard et
252 al., 2003) to assess stereochemical soundness, allowing selection of the final best
253 model. VGSC mutations were introduced into the homology model, followed by 30
254 steps of steepest descent and then 50 steps of conjugate gradient energy minimization,
255 using Swiss-PdbViewer software (Guex et al., 1999).

256 A 3-dimensional structure file for fenpropathrin, permethrin or beta-
257 cypermethrin was generated using MarvinSketch (v19.22) of the ChemAxon suite
258 (<http://www.chemaxon.com>). These files were then prepared using the
259 AutoDockTools program (v.1.5.7; Molecular Graphics Laboratory, Scripps Research
260 Institute, La Jolla, CA, USA) to define rotatable bonds and merge nonpolar
261 hydrogens. The Autodock Vina (Trott & Olson, 2010) program was used for docking
262 the ligands within a grid of 20 x 20 x 20 points of 1 Å spacing that was centered on

263 the PyR1 binding pocket (O'Reilly et al., 2006a, Du et al., 2015) of the WT VGSC
264 model. Figures were produced using Pymol 2.2.3 (DeLano Scientific, San Carlos, CA,
265 USA).

266 **Statistical analyses**

267 PoloPlus v.2.0 (LeOra Software 2008) was used to calculate the median lethal
268 concentration (LC₅₀) and 95% confidence limits (95% CL) for bioassays of *P. citri*
269 and transgenic flies. Student's *t*-tests were used to test differences between
270 experimental and control groups and also between mutant and WT groups. The
271 GraphPad Prism 8.0 software program (GraphPad Software Inc.) was used for data
272 analysis and plot generation.

273 **Results**

274 **Resistance to fenprothrin**

275 The fenprothrin resistance of *P. citri* collected from four field sampling sites
276 varied (Table 1). The LC₅₀ values of fenprothrin against the *P. citri* field
277 populations from Yuxi (YX-YN), Ganzhou (GZ-JX), Liangping (LP-CQ), Anyue (AY-
278 SC), and Nanning (NN-GX) ranged from 45.5 to 4,093.9 mg/L in 2021. The WZ-CQ
279 population was more susceptible to fenprothrin than the NN-GX, Tongnan (TN-
280 CQ), Wanzhou (WZ-CQ) and Guilin (GL-GX) populations collected in 2022. Except
281 for the LP-CQ, TN-CQ, and WZ-CQ populations, very high levels of resistance
282 (i.e., >3,000-fold resistance levels) were detected in field populations compared to the
283 SS strain.

284 **VGSC mutations in *P. citri* populations**

285 To detect potential VGSC mutations in *P. citri* populations of southwest China,
286 the presence of two potential mutation sites (L1031 and F1751) of *P. citri* VGSC gene
287 (the amino acid positions are numbered based on the *P. citri* VGSC protein sequence)
288 were investigated that corresponded to the L1024 and F1538 mutations of *T. urticae*
289 (Inak et al., 2019). Three mutation types were observed for *P. citri*, with the L1031V
290 amino acid change at segment 6 of domain II (DII S6) in addition to the F1747L and
291 F1751I mutations at segment 6 of domain III (DIII S6) (Fig. 1) in the resistant
292 populations of YX-YN, AY-SC, NN-GX, and GL-GX.

293 **Frequency of the L1031V, F1747L, and F1751I mutations in field populations**

294 The mutations in over 14,000 mites were evaluated using amplicon sequencing
295 of genes encoded by individuals from nineteen different sites across four provinces
296 (Table 2). The frequency of mutations varied from 11.6 to 82.1%, 0 to 34.8%, and 0.5
297 to 31.8% for L1031V, F1747L and F1751I, respectively. The L1031V mutation was
298 widespread across sample sites, with highest frequency of 82.1% in the GL2-GX
299 population. The F1751I mutation was most prevalent at the Liangping locality in
300 2022, occurring in 31.8% of genotypes. Moreover, the F1751I mutation was
301 widespread among the nineteen samples. A new mutation (F1747L) that had not
302 previously been reported in pest mites was also identified in twelve samples, with the
303 highest frequency (34.8%) observed in the RL-YN samples.

304 **Correlation of fenpropathrin resistance level and VGSC mutation frequency in *P.***
305 ***citri* field populations**

306 When the mutation frequencies were plotted against the fenpropathrin resistance
307 levels in *P. citri*, the L1031V, F1747L, and F1751I mutations were lowly correlated
308 with the fenpropathrin resistance level ($r^2 = 0.1166, 0.1262$ and 0.0092 , respectively)
309 (Fig. S2). In addition, low correlations were also observed between the frequencies of
310 mutation combinations and the resistance levels ($r^2 = 0.0649, 0.1052$ and 0.0632 for
311 L1031V+F1747L, L1031V+F1751I and L1031V+F1747L+F1751I, respectively)
312 (Fig. S2). Nevertheless, the upward tendency of resistance ratio was recorded when
313 the allele frequency increased.

314 **Importance of the VGSC mutation in pyrethroid resistance using transgenic**
315 **expression in *Drosophila melanogaster***

316 Bioassays with fenpropathrin revealed that flies with the VGSC mutations
317 L1031V, F1747L, and F1751I were resistant to fenpropathrin (Table 3). However,
318 different contributions of VGSC mutations to fenpropathrin resistance were observed
319 for transgenic flies. Varying LC_{50} values of 4.17, 2.97, and 5.63 mg/L were identified
320 in the Gal4/UAS-mutant (L1031V, F1747L, and F1751I) strains. Compared to the
321 Gal4/UAS-WT strain, the resistance of the Gal4/UAS-mutant (L1031V, F1747L, and
322 F1751I) strains were significantly higher (2.71-, 1.93-, and 3.66-fold higher,

323 respectively), with no overlap at the 95% CL level. Combinations of VGSC mutations
324 in transgenic flies further increased resistance to fenpropathrin compared to single
325 mutations (Table 3). Indeed, the highest resistance (4.55-fold compared to the
326 Gal4/UAS-WT lines) was detected when all three sites were mutated. Similar results
327 were observed in the UAS-VGSC lines (Table S6).

328 Bioassays with permethrin (type I pyrethroid) and beta-cypermethrin (type II
329 pyrethroid) were also performed to assess their contributions to pyrethroid resistance
330 in transgenic lines (Fig. 2). Flies with VGSC mutations exhibited lower mortality and
331 knockdown rates compared to WT flies. The Gal4/UAS-mutant (L1031V, F1747L,
332 F1751I, L1031V+F1747L, L1031V+F1751L, and L1031V+F1747L+F1751I) lines
333 exhibited significantly lower mortality and knockdown rates (76, 81, 73.6, 64, 51, and
334 50%, respectively) to permethrin compared to the Gal4/UAS-WT line (94%) at 48 h
335 ($p < 0.001$) (Fig. 2). No significant differences were observed between flies with
336 overexpressed VGSC genes (Gal4/UAS lines) compared to control (UAS line) flies
337 (Fig. S3). Similar results were observed with beta-cypermethrin bioassays, with
338 significantly reduced mortalities and knockdown rates in Gal4/UAS-mutant lines (70,
339 79, 70.4, 68, 60, and 56%, respectively) compared with Gal4/UAS-WT flies (90%)
340 after exposure for 48 h ($p < 0.001$) (Fig. 2). For Gal4/UAS and UAS lines exposed to
341 beta-cypermethrin, flies with VGSC-WT and VGSC-L1031V+F1747L genes
342 exhibited significant differences in mortalities and knockdowns at exposure times of
343 24 and 36 h (Fig. S4).

344 These results indicate that mutations of *P. citri* VGSC genes confer resistance to
345 pyrethroids, but with varying degrees of resistance. The F1751I mutation conferred
346 the highest resistance to pyrethroids, followed by the L1031V and F1747L mutations.

347 **Molecular modeling of the VGSC mutations**

348 A homology model of *P. citri* VGSC was generated in order to assess how
349 mutations could affect pyrethroid binding. Although L1031 is located on the DII S6
350 while F1747 and F1751 are located on the DIII S6 (Fig. 1A), in the 3-dimensional
351 model (Fig. 3A, B), these three residues are positioned in close spatial proximity with
352 $<13 \text{ \AA}$ distance between their α -carbons. Their side chains shape the lower end of the

353 lipid-accessible cavity that forms pyrethroid binding site PyR1 (O'Reilly et al., 2006a,
354 Du et al., 2015).

355 A docking prediction for permethrin with an estimated free energy of binding
356 (ΔG_b) of -8.15 kcal/mol positioned the pyrethroid to occupy the length of the binding
357 cavity (Fig. 3A and Fig. S5B, E). The phenoxy group of permethrin makes direct
358 binding interactions with F1747 and F1751 side chains, suggesting stacking between
359 the three aromatic rings is a major binding interaction. In contrast, the L1031 side-
360 chain is too far ($>4.5 \text{ \AA}$) to directly interact with permethrin but it does contact the
361 F1751 side chain and pack against the DIII S6 helix, thus contributing to shaping the
362 binding site. Similar docking poses, estimated binding energies and interactions with
363 F1747 and F1751 side chains were found with fenpropathrin (ΔG_b of -8.0 kcal/mol)
364 and beta-cypermethrin (ΔG_b) of -6.78 kcal/mol (Fig. S5).

365 The loss of an aromatic ring with either the F1747 or F1751 mutation could
366 reduce aromatic-aromatic binding interactions with a pyrethroid phenoxy group,
367 which is common to permethrin, fenpropathrin and beta-cypermethrin. Furthermore,
368 F1747L and F1751I mutations each introduce a branched hydrocarbon side-chain that
369 reshapes the binding cavity in that region (Fig. 3B-E), which may sterically hinder
370 pyrethroid binding. In contrast, L1031V is not predicted to affect pyrethroid binding
371 directly but the mutation to a smaller side chain may affect packing between DII and
372 DIII S6 helices, thereby conferring resistance by allosterically modifying the
373 pyrethroid binding site.

374 **Discussion**

375 In this study, different degrees of resistance to the pyrethroid fenpropathrin were
376 identified among mites in most citrus orchards of Southern China. The most common
377 mechanism of pyrethroid resistance has arisen from substitution of certain amino
378 acids in pest VGSCs (Dong et al., 2014). Indeed, many such mutations associated
379 with pyrethroid resistance have been described in diverse arthropod species, including
380 substitution of M918T/L/V/I, L925I/V, T929I/C/V/N, L932F, L1014F/H/S/W,
381 F1534C, and F1538I (numbering based on *Musca domestica* VGSC) (Scott, 2019,
382 Dong et al., 2014). Limited mutations of VGSCs have been identified in pest mite

383 populations with pyrethroid resistance, with the few being L925V/I (Koc et al., 2021,
384 Almecija et al., 2022), L1024V, A1215D, and F1538I (numbering according to *M.*
385 *domestica*) (Simma et al., 2020, Zhang et al., 2022, Ding et al., 2015). The L1031V
386 (L1024), F1747L (F1534), and F1751I (F1538) mutations were detected in field-
387 collected populations of *P. citri* in this study. However, the L925 and A1215 mutations
388 of VGSCs were not identified in these *P. citri* populations. In addition, the F1747L
389 (F1534) mutation identified in pest mites of this study represent the first identification
390 of this mutation in mites. The F1747L (F1534) mutation of the IIS6 site of VGSC
391 was also identified in DDT/permethrin-resistant *Aedes aegypti* and was closely
392 associated with a resistant phenotype (Yanola et al., 2011).

393 The frequencies of point mutations in *P. citri* that were related to pyrethroid
394 resistance were evaluated with amplicon sequencing, as previously conducted for *H.*
395 *armigera* (Jin et al., 2018). The L1031V and F1751I mutations exhibited frequencies
396 of 11.6–82.1% and 0.5–31.8%, respectively, in field populations collected from
397 nineteen citrus producing sites, while the F1747L mutation occurred in ten field sites
398 and had a frequency of 1.1–34.8%. The frequency of F1538I in VGSCs of *T. urticae*
399 field populations ranged from 10–100% (Zhang et al., 2022). The high frequency of
400 the F1747I (F1538I) mutation observed in the current study is consistent with other
401 studies of *P. citri* (Ding et al., 2015), *T. urticae* (Riga et al., 2017), *Panonychus ulmi*
402 (Rameshgar et al., 2019), *Dermanyssus gallinae* (Katsavou et al., 2020), and
403 *Tetranychus cinnabarinus* (Xu et al., 2013). The L1031V (L1024) mutation has also
404 been detected in *T. urticae* (Kwon et al., 2010, Simma et al., 2020) and *P. ulmi*
405 (Rameshgar et al., 2019) field populations, where it conferred pyrethroid resistance.
406 When the relations were analyzed between the frequencies of each mutation (and their
407 combinations) and the resistance level in field populations, no positive relations were
408 observed between the VGSC mutation frequency and the resistance levels of *P. citri*
409 field populations to fenpropathrin. The diverse acaricides application usually led to
410 the complex resistance evolution within a complex genetic background in mites. This
411 phenomenon was more likely caused by some factors, for instance, the increased
412 detoxification enzymes activities, cross-resistance occurrence. Previous studies

413 indicated that carboxylesterases genes were directly involved in fenprothrin
414 resistance of the spider mites, *Tetranychus cinnabarinus* (Boisduval) (Wei et al.,
415 2019) and *P. citri* (Shen et al., 2016). What's more, glutathione S-transferase and
416 cytochrome P450 monooxygenases might contribute to fenprothrin resistance in
417 mites as well (Shen et al., 2014, Liao et al., 2018, Shi et al., 2016). The bioassay
418 showed that moderate cross-resistance to fenprothrin occurred when *Tetranychus*
419 *urticae* (Koch) developed extremely resistance to fenpyroximate (Kim et al., 2004).
420 Similarly, a significant reduced susceptibility to fenprothrin was observed after an
421 extremely abamectin resistance was developed in *T. urticae* (Sato et al., 2005). More
422 importantly, resistance monitoring indicated that field populations of *P. citri* had
423 developed different degrees of resistance to all tested acaricides (Pan et al., 2020). All
424 these potential factors led to the current negative correlation between fenprothrin
425 resistance and VGSC mutation frequency in *P. citri*. Thus, more evidences need to be
426 supplied by other functional verification, like the bioassay in transgenic flies after
427 VGSC mutation introduction.

428 Therefore, we introduced the VGSC mutations L1031V, F1747L, and F1751I
429 into the model organism *Drosophila melanogaster* through CRISPR/Cas9 genome
430 editing in order to functionally validate the ability of these mutations to confer
431 insecticide resistance *in vivo*. Bioassays revealed significantly increased resistance (in
432 the order of F1751I > L1031V > F1747L) to pyrethroids (both types I and II) in
433 transgenic flies carrying VGSC mutations compared to control flies (VGSC-WT). The
434 F1747L (F1534L) mutation conferred lower resistance to types I and II pyrethroids. In
435 contrast to previous observation, the F1534L mutation did not affect channel
436 resistance to pyrethroids in *Aedes albopictus* (Gao et al., 2018, Xu et al., 2016). In a
437 previous study, the F1534L mutation significantly reduced channel sensitivity to Type
438 I pyrethroids, but not to two Type II pyrethroids (Yan et al., 2020), contrasting with
439 our results. It is possible that different insect genomic backgrounds may underly the
440 differences in sensitivity in bioassays among different studies, possibly via the
441 presence of additional mutations or nucleotide sequence polymorphisms (Yan et al.,
442 2020). The relationships between VGSC mutations and pyrethroid resistance

443 consequently require further studies, such as cryo-electron microscopy investigation.

444 *In vivo* functional validation has been investigated for L1024V, revealing channel
445 sensitivity in oocytes and changes in channel sensitivity to pyrethroids (Du et al.,
446 2015). A previous study also observed that the F1538I mutation occurred in the WZ
447 population of *P. citri* (Ding et al., 2015). Previous identification of mutations in
448 VGSCs have suggested that F1538I conferred strong resistance to pyrethroids in *T.*
449 *urticae* (Tsagkarakou et al., 2009) and *T. cinnabarinus* (Feng et al., 2011). The
450 bioassays in this study further confirmed that the F1538I mutation of VGSCs plays an
451 important role in the strong resistance to pyrethroids. Combinations of VGSC
452 mutations (e.g., L1031V+F1751I and L1031V+F1747L+F1751I) were also observed
453 in several *P. citri* field populations in this study. The L1031V+F1747L+F1751I
454 mutation combination (equivalent to L1024V+F1534L+F1538I) was first detected in
455 highly fenprothrin-resistant populations of *P. citri*. A similar combination
456 (L1024V+F1538I) was also detected in resistant *P. ulmi* populations and was shown
457 to confer pyrethroid resistance (Rameshgar et al., 2019). An additional mutation
458 combination (A1215D+F1538I and L1024V+A1215D) was identified in *T. urticae*
459 VGSCs among different resistant strains or populations (Kwon et al., 2010, Xu et al.,
460 2018). When these mutation combinations (L1031V+F1751I and
461 L1031V+F1747L+F1751I) were introduced to transgenic flies, significant pyrethroid
462 resistance was observed compared to control flies. Thus, mutations in combination
463 can further increase resistance to pyrethroids. After these mutations were introduced
464 into the *Drosophila*, although a small change in resistance was recorded, with the
465 highest resistance of only 4.55-fold, it is reliable based on a recent report in *T. urticae*.
466 Selection with abamectin for generations resulted in a resistance ratio of 105 in the
467 mite. Subsequently, the introduction of glutamate-gated chloride channel I321T into
468 transgenic flies **through CRISPR/Cas9 genome editing** led to only a 2.66-fold
469 resistance to abamectin (Xue et al., 2021).

470 Our modelling indicates that all three resistance-associated mutations are brought
471 into close proximity in the folded protein in a region of PyR1 that is also lined by the
472 DII S4-S5 linker and DII S5 (O'Reilly et al., 2006b). Our previous electrophysiology

473 study demonstrated that resistance-associated mutations in this region greatly
474 diminish the effect of pyrethroids with a phenoxy group have but not fenfluthrin,
475 which lacks the terminal phenyl ring, nor DDT (Usherwood et al., 2007). Numerous
476 modelling studies have predicted that a pyrethroid's phenoxy group binds in this
477 lower portion of PyR1 (O'Reilly et al., 2006b, Usherwood et al., 2007, O'Reilly et al.,
478 2014, Wu et al., 2017) and here we propose aromatic-aromatic interactions between
479 F1747 and F1751 side-chains and a pyrethroid's terminal phenyl ring as a specific
480 binding interaction. If F1747L and F1751I mutations confer resistance through our
481 proposed steric hinderance mechanism with a pyrethroid's phenoxy group, then use
482 of pyrethroids lacking a terminal phenyl ring may overcome this resistance.
483 Alternatively, our previous modelling predicts that miticidal analogues of DDT bind
484 in the upper portion of PyR1 (O'Reilly et al., 2014) and so may be too distant from
485 F1747 and F1751 to be affected by their mutations. Assessing alternatives to the
486 currently-used pyrethroids to overcome resistance in mutant *P.citri* populations will
487 therefore require field trials and further experimentation.

488 In conclusion, high resistance to fenpropathrin was detected in populations of *P.*
489 *citri* collected from the field. Amplicon sequencing revealed the frequencies of the
490 mutations L1031V, F1747L and F1751I of the *P. citri* VGSC gene, with L1031V and
491 F1751I occurring in all evaluated populations at frequencies of 11.6–82.1% and 0.5–
492 31.8%, respectively. The action of pyrethroids (types I and II) was also demonstrated
493 in transgenic flies. Compared to VGSC-WT flies, significant decreases in sensitivity
494 were observed in VGSC-mutant flies to varying degrees. Modeling of the *P. citri*
495 VGSC suggested that pyrethroids are located at PyR1 and that interactions were
496 disturbed after the substitution of the residues at the L1031, F1747, and F1751 sites.
497 Consequently, the interaction to pyrethroids in *P. citri* was directly caused by the
498 substitution of certain residues that affect interactions with pyrethroids.

499 **Acknowledgement**

500 We thank Menghao Xia and Yuchuang Li for helping to collect mites. This study was
501 supported by the Science and Technology Basic Resources Investigation Program of
502 China (2018FY101105), the National Natural Science Foundation of China

503 (31972272, 31871969), the Fundamental Research Funds for the Central Universities
504 (SWU-XDPY22001) of China, and the China Agricultural Research System of MOA
505 and MARA.

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717

Figure legends

Fig. 1. A. Schematic showing the *P. citri* voltage gated sodium channel (VGSC). Alignment of amino acid sequences in the IIS6 (B) and IIIS6 (C) regions indicating the alignment positions of the mutations L1031V, F1747L, and F1751I.

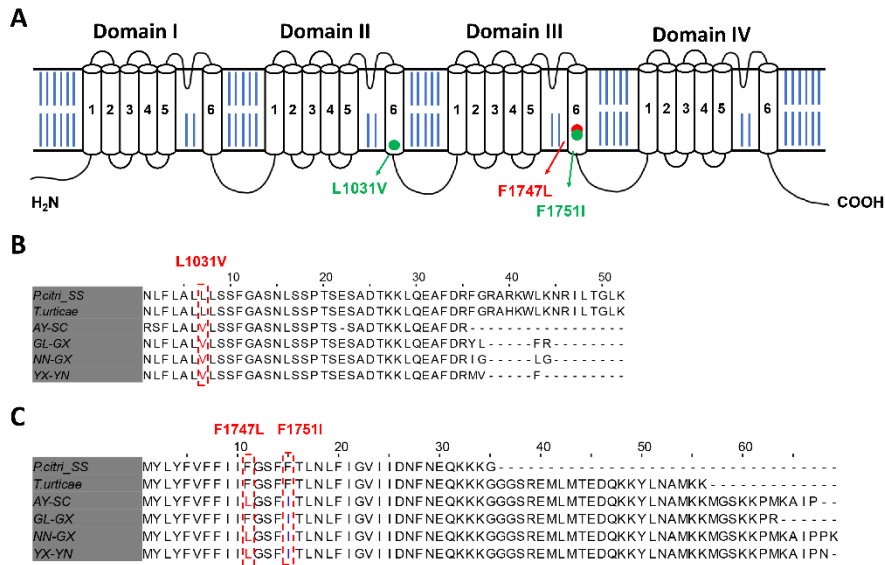


Fig. 2. Pyrethroid resistance bioassays of transgenic flies. Data are means \pm S.E.M. Statistically significantly differences are indicated as: ***: $p < 0.001$. “*” was only showed at most close to WT-lines for 48 h and the rest were omitted with a significant difference too.

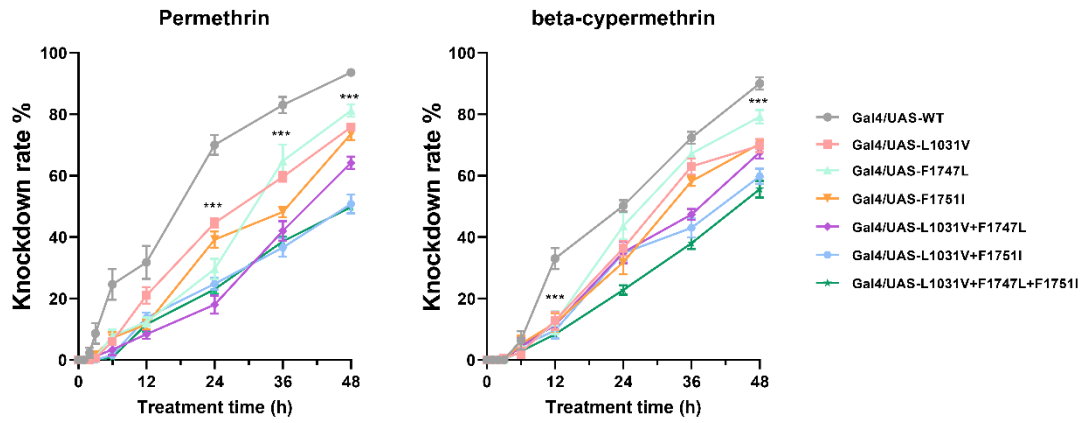


Fig. 3. Homology model of the *P. citri* sodium channel showing residues at position 1031 (brown), 1747 (green) and 1751 (orange) (A) A docking prediction for permethrin shown in yellow space-fill within the lipid-accessible pyrethroid binding site. (B-C) The wildtype channel is shown in ribbon (B) or surface representation (C). (D-E) the triple-mutant channel is shown in ribbon (D) or surface representation (E).

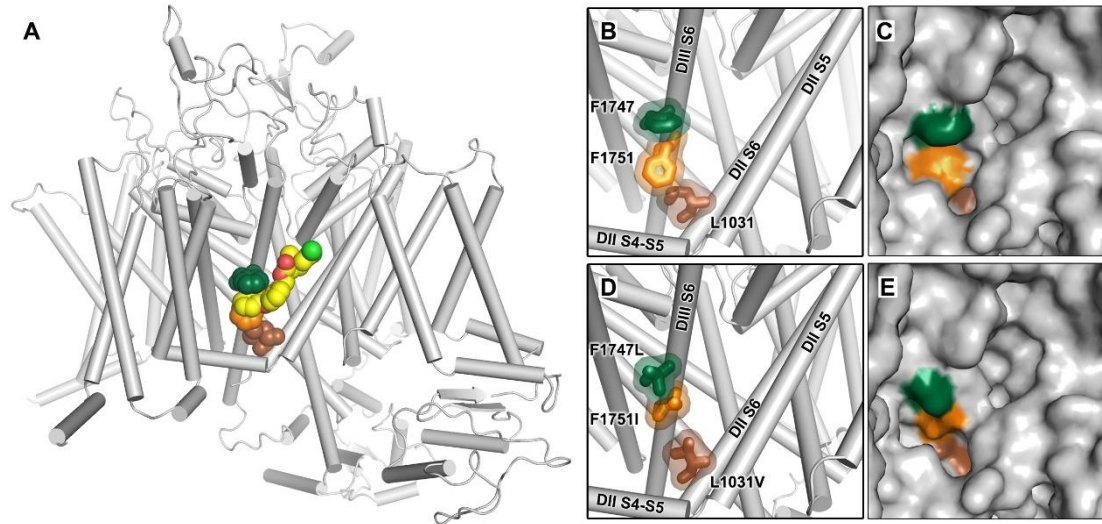


Table 1. Toxicity of fenpropathrin in eight field populations of *Panonychus citri* sampled from 2021 to 2022.

Population	Year	N	Slope (\pm SE)	LC ₅₀ (mg/L)	95% CL ^a	χ^2 (df)	RR ^b
SS	-	481	1.026 \pm 0.011	0.266	0.196-0.371	4.371 (3)	1
YX-YN	2021	595	1.502 \pm 0.119	1300.287	989.122-1795.749	6.8534 (4)	4888.30
GZ-JX	2021	932	2.851 \pm 0.157	854.885	541.823-1328.939	14.590 (5)	3213.85
LP-CQ	2021	635	3.107 \pm 0.217	45.468	29.600-65.761	14.490 (3)	170.93
AY-SC	2021	578	4.380 \pm 0.558	4093.913	3690.024-4496.152	1.700 (5)	15390.65
NN-GX	2021	521	2.253 \pm 0.194	1668.069	1021.821-2894.240	10.396 (3)	6270.93
NN-GX	2022	540	3.498 \pm 0.443	2771.812	29.600-65.761	1.9 (4)	10420.35
TN-CQ	2022	568	1.242 \pm 0.117	191.580	155.559-240.830	3.757 (3)	760.23
WZ-CQ	2022	550	1.107 \pm 0.150	3.217	2.562-4.249	1.760 (3)	12.09
GL-GX	2022	480	2.452 \pm 0.211	1395.170	2470.044-3046.003	4.7919 (3)	5245.00

^aCL, confidence limits.

^bRR, resistance ratio = LC₅₀ of the field population / LC₅₀ of the susceptible strain.

YX-YN, collected from Yuxi, Yunnan province (E103°7'20", N24°12'15"); GZ-JX: Ganzhou, Jiangxi province (E114°52'4.44", N25°47'21.48"); LP-CQ; Liangping, Chongqing (E107°39'37.02", N30°36'48.70"); AY-SC: Anyue, Sichuan province (E105°18'36", N30°8'32"); NN-GX: Nanning, Guangxi province (E108°31'48", N23°14'49.2"); TN-CQ: Tongnan, Chongqing city (E105°46'2", N30°4'47"); WZ-CQ: Wanzhou, Chongqing city (E108°30'4", N30°56'3"); GL-GX: Guilin, Guangxi province (E110°17'14", N25°28'33").

Table 2. Frequency of the L1031V, F1747L and F1751I mutations in in VGSCs sequenced from pools of field-captured *P. citri* from southwest China.

Population	Date	Mutation site and frequency (Mean \pm SEM) (%)		
		L1031V	F1747L	F1751I
TN1-CQ	2021, 3	43.8 \pm 0.9	11.3 \pm 0.9	13.2 \pm 0.4
TN2-CQ	2022, 4	51.0 \pm 1.7	10.6 \pm 0.4	14.6 \pm 1.2
LP1-CQ	2021, 4	44.8 \pm 1.4	13.3 \pm 0.5	31.8 \pm 0.4
LP2-CQ	2022, 6	68.7 \pm 1.1	0	14.1 \pm 0.4
WZ-CQ	2021, 4	26.7 \pm 1.4	1.20 \pm 0.1	0.50 \pm 0.1
LZ-SC	2021, 3	45.6 \pm 0.5	0	19.6 \pm 0.7
ZZ-SC	2021, 4	46.1 \pm 0.7	0	17.8 \pm 0.6
YB-SC	2021, 6	40.0 \pm 1.9	0	21.0 \pm 0.2
NJ-SC	2021, 5	41.3 \pm 1.7	33.7 \pm 0.5	13.2 \pm 0.4
LS-SC	2021, 5	45.6 \pm 0.2	15.2 \pm 1.2	20.8 \pm 0.4
AY-SC	2021, 7	49.6 \pm 0.5	0	19.5 \pm 0.4
NN1-GX	2021, 9	40.9 \pm 1.8	15.7 \pm 0.3	27.8 \pm 1.2
NN2-GX	2022, 7	74.6 \pm 0.4	1.10 \pm 0.2	25.8 \pm 0.7
GL1-GX	2021, 5	48.1 \pm 1.3	0	18.1 \pm 0.3
GL2-GX	2022, 7	82.1 \pm 0.3	1.50 \pm 0.2	16.4 \pm 0.6
YX-YN	2021, 5	11.6 \pm 0.7	19.2 \pm 0.9	4.60 \pm 0.3
WSZ-YN	2021, 10	43.5 \pm 0.8	7.70 \pm 1.0	14.7 \pm 0.1
GZ-JX	2021, 5	24.4 \pm 0.10	3.4 \pm 0.26	45.3 \pm 0.42
RL-YN	2021, 11	27.4 \pm 0.4	34.8 \pm 1.3	19.3 \pm 0.3

TN1/2-CQ, collected from Tongnan, Chongqing city (E105°45'11", N30°9'9"; E105°46'2", N30°4'47"); LP1/2-CQ: Liangping, Chongqing city (E107°39'37.02", N30°36'48.70"; E107°42'6", N30°36'47"); WZ-CQ: Wangzhou, Chongqing city (E108°41'49.92", N28°53'56.4"); LZ-SC: Luzhou, Sichuan province (E105°34'8.4", N28°53'56.4"); ZZ-SC: Zizhong, Sichuan province (E104°54'58", N29°41'31"); YB-SC: Yibing, Sichuan province (E104°25'27", N28°37'54"); NJ-SC: Neijiang, Sichuan province (N29°2'40", E105°1'1"); LS-SC: Leshang, Sichuan province (N29°2'40", E103°34'51.80"); AY-SC: Anyue, Sichuan province (E105°18'36", N30°8'32"); NN1/2-GX: Nanning, Guangxi province (E108°31'48", N23°14'49.2"); GL1/2-GX: Guilin, Guangxi province (E110°27'25.78", N25°19'20.28"; E110°17'14", N25°28'33"); YX-YN: Yuxi, Yunnan province (E103°7'20", N24°12'15"); WSZ-YN: Wenshanzhou, Yunnan province (E103°56'22", N23°10'15"); GZ-JX: Ganzhou, Jiangxi province (E114°52'4.44", N25°47'21.48"); RL-YN: Ruili, Yunnan province (E97°52'12", N24°1'9").

Table 3. Bioassays of fenpropathrin resistance in transgenic flies.

Strains	N	Slope (\pm SE)	LC ₅₀ (95%CL) ^a (mg/L)	χ^2 (df)	RR ^b
Gal4/UAS -WT	286	1.272 \pm 0.243	1.54 (1.06-2.31)	0.143 (4)	1
Gal4/UAS - L1031V	270	1.689 \pm 0.234	4.17 (3.35-5.31)	0.417 (3)	2.71
Gal4/UAS - F1747L	289	2.440 \pm 0.289	2.97 (2.47-3.56)	0.102 (3)	1.93
Gal4/UAS - F1751I	278	2.125 \pm 0.403	5.63 (4.57-7.11)	0.175 (4)	3.66
Gal4/UAS - L1031V+ F1747L	282	0.991 \pm 0.171	4.73 (3.17-8.22)	1.774 (3)	3.08
Gal4/UAS - L1031V+ F1751I	288	1.526 \pm 0.244	5.82 (4.51-8.20)	1.373 (4)	3.78
Gal4/UAS - L1031V+ F1747L+ F1751I	288	2.637 \pm 0.295	6.99 (4.98-8.15)	1.232 (2)	4.55

^aCL, confidence limits.

^bRR, resistance ratio = LC₅₀ of the mutant lines / LC₅₀ of the WT lines.

Appendix A. Supplementary material

Table S1. Locations, origins, and geographic information for *Panonychus citri* populations collected for bioassays.

Population	Location	Collection date	Host plant	Longitude and latitude	
YX-YN	Yuxi, Yunan	2021, 5	<i>Citrus reticulata</i> 'Unshiu'	E103°7'20"	N24°12'15"
GZ-JX	Ganzhou, Jiangxi	2021, 5	<i>Citrus limon</i>	E114°52'4.44"	N25°47'21.48"
LP-CQ	Liangping, Chongqing	2021, 6	<i>Citrus maxima</i> (Burm.) Merr.cv. <i>Liangping Yu</i>	E107°39'37.02"	N30°36'48.70"
AY-SC	Anyue, Sichuan	2021, 7	<i>Citrus limon</i>	E105°18'36"	N30°8'32"
NN-GX	Nanning, Guangxi	2021, 9	<i>Fertile orange</i>	E108°31'48"	N23°14'49.2"
		2022, 7			
TN-CQ	Tongnan, Chongqing	2022, 4	<i>Citrus limon</i>	E105°46'2"	N30°4'47"
WZ-CQ	Wanzhou, Chongqing	2022, 6	<i>Citrus sinensis</i>	E108°30'4"	N30°56'3"
GL-GX	Guilin, Guangxi	2022, 7	<i>Citrus reticulata</i>	E110°17'14"	N25°28'33"

Table S2. Locations, origins, and geographic information for *Panonychus citri* populations collected for mutation frequency analysis.

Population	Collection location	Collection date	Host plant	Longitude and latitude	
TN1-CQ	Tongnan, Chongqing	2021, 3	<i>Citrus limon</i> (L.) Burm. F.	E105°45'11"	N30°9'9"
TN2-CQ	Tongnan, Chongqing	2022, 4	<i>Citrus limon</i> (L.) Burm. F.	E105°46'2"	N30°4'47"
LP1-CQ	Liangping, Chongqing	2021, 4	<i>Citrus maxima</i> (Burm.) Merr.cv. Liangping Yu	E107°39'37.02"	N30°36'48.70"
LP2-CQ	Liangping, Chongqing	2022, 6	<i>Citrus reticulata</i> Banco	E107°42'6"	N30°36'47"
WZ-CQ	Wangzhou, Chongqing	2021, 4	<i>Citrus reticulata</i> 'Banco'	E108°41'49.92"	N28°53'56.4"
LZ-SC	Luzhou, Sichuan	2021, 3	<i>Citrus sinensis</i>	E105°34'8.4"	N28°53'56.4"
ZZ-SC	Zizhong, Sichuan	2021, 4	<i>Citrus sinensis</i>	E104°54'58"	N29°41'31"
YB-SC	Yibing, Sichuan	2021, 6	<i>Citrus reticulata</i> 'Ponkan	E104°25'27"	N28°37'54"
NJ-SC	Neijiang, Sichuan	2021, 5	<i>Citrus kanper</i>	N29°2'40"	E105°1'1"
LS-SC	Leshang, Sichuan	2021, 5	<i>Citrus reticulata</i> 'Banco'	N29°2'40"	E103°34'51.80"
AY-SC	Anyue, Sichuan	2021, 7	<i>Citrus limon</i> (L.) Burm. F.	E105°18'36"	N30°8'32"
NN1-GX	Nanning, Guangxi	2021, 9	Fertile orange	E108°31'48"	N23°14'49.2"
NN2-GX	Nanning, Guangxi	2022, 7	Fertile orange	E108°31'48"	N23°14'49.2"
GL1-GX	Guilin, Guangxi	2021, 5	<i>Citrus reticulata</i> 'Banco'	E110°27'25.78"	N25°19'20.28"
GL2-GX	Guilin, Guangxi	2022, 7	<i>Citrus reticulata</i> 'Banco'	E110°17'14"	N25°28'33"
YX-YN	Yuxi, yunnan	2021, 5	<i>Citrus reticulata</i> 'Unshiu'	E103°7'20"	N24°12'15"
WS-YN	Wenshanzhou, yunnan	2021, 10	Tribute Citru	E103°56'22"	N23°10'15"
GZ-JX	Ganzhou, Jiangxi	2021, 5	<i>Citrus limon</i> (L.) Burm. F.	E114°52'4.44"	N25°47'21.48"
RL-YN	Ruili, yunnan	2021, 11	<i>Citrus limon</i> (L.) Burm. F.	E97°52'12"	N24°1'9"

Table S3. PCR primers used to amplify VGSC fragments and detect mutants.

Primer name	Direction	Sequence (5'-3')	Products (bp)
<i>Pc</i> VGSC_L1031V	Forward	CGTGTTCTTTGIGGTGAATG	236
	Reverse	TTCTGAGCTCTACCGAAACG	
<i>Pc</i> VGSC_F1747L	Forward	GCAACATTCAAGGGTTGGAC	271
	Reverse	GGAATCGCTTTCATCGGCTT	
<i>Pc</i> VGSC_F1751I	Forward	GCAACATTCAAGGGTTGGAC	271
	Reverse	GGAATCGCTTTCATCGGCTT	

Table S4. Barcodes used in PCR amplification of VGSCs of *P. citri* populations to enable multiplex sequencing.

Population	Primer tag
TN1-CQ	CGATGT
TN2-CQ	
LP1-CQ	GGCTAC
LP2-CQ	
WZ-CQ	ATCACG
LZ-SC	TTAGGC
ZZ-SC	TGACCA
YB-SC	CAGATC
NJ-SC	ACTTGA
LS-SC	GATCAG
AY-SC	TGACCA
NN1-GX	ACAGTG
NN2-GX	
GL1-GX	GGCTAC
GL2-GX	
YX-YN	CGATGT
WSZ-YN	TTAGGC
RL-YN	ACAGTG

Table S5. PCR primers used to validate VGSC gene integration in transgenic fly constructs.

Transgenic lines	Direction	Sequence (5'-3')
UAS-VGSC-pm-WT	Forward	CAAGACTATCTGTGATCAAC
	P10-R	GCCACTAGCTCGCTATACT
UAS-VGSC-pm-L1031V	Forward	AGAGATACCCCGTTGGAATTT
	P10-R	GCCACTAGCTCGCTATACT
UAS-VGSC-pm-F1747L	Forward	CAAGACTATCTGTGATCAAC
	P10-R	GCCACTAGCTCGCTATACT
UAS-VGSC-pm-F1751I	Forward	AGAGATACCCCGTTGGAATTT
	P10-R	GCCACTAGCTCGCTATACT
UAS-VGSC-pm- L1031V+ F1747L	Forward	AGAGATACCCCGTTGGAATTT
	P10-R	GCCACTAGCTCGCTATACT
UAS-VGSC-pm- L1031V+ F1751I	Forward	AGAGATACCCCGTTGGAATTT
	P10-R	GCCACTAGCTCGCTATACT
UAS-VGSC-pm- L1031V+ F1747L+ F1751I	Forward	AGAGATACCCCGTTGGAATTT
	P10-R	GCCACTAGCTCGCTATACT

Table S6. Bioassays of transgenic fly resistance to pyrethroids.

Strains	N	Slope (\pm SE)	LC ₅₀ (95%CI) ^a (mg/L)	χ^2	RR ^b
UAS - WT	280	1.823 \pm 0.271	1.64 (1.25-2.24)	0.080	1
UAS - L1031V	265	1.817 \pm 0.235	3.42 (2.75-4.23)	0.415	2.08
UAS - F1747L	276	2.052 \pm 0.283	3.60 (2.78-4.59)	0.503	2.19
UAS - F1751I	287	1.833 \pm 0.251	4.45 (3.59-5.67)	0.268	2.71
UAS - L1031V+ F1747L	290	1.109 \pm 0.179	5.81 (4.01-9.72)	1.267	3.53
UAS - L1031V+ F1751I	287	1.776 \pm 0.253	5.26 (4.21-6.91)	1.673	3.20
UAS - L1031V+ F1747L+ F1751I	270	2.128 \pm 0.386	5.73 (4.68-7.18)	0.282	3.48

Fig. S1. Alignment of amino acid sequences in the IIS6 (A) and IIS6 (B) regions of VGSCs to verify the mutations L1031V, F1747L, and F1751I in transgenic flies.

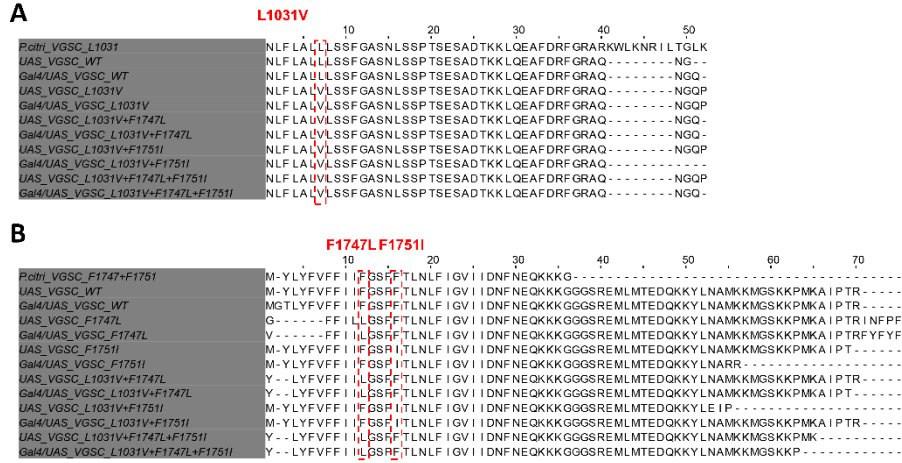


Fig. S2. Correlation between the fenpropathrin resistance level and the frequency of VGSC mutations in the *P. citri* field populations. Resistance ratios of field populations to fenpropathrin were plotted against the frequency of VGSC mutations with single or combinations. Regression equation and correlation coefficient were included on the graphs.

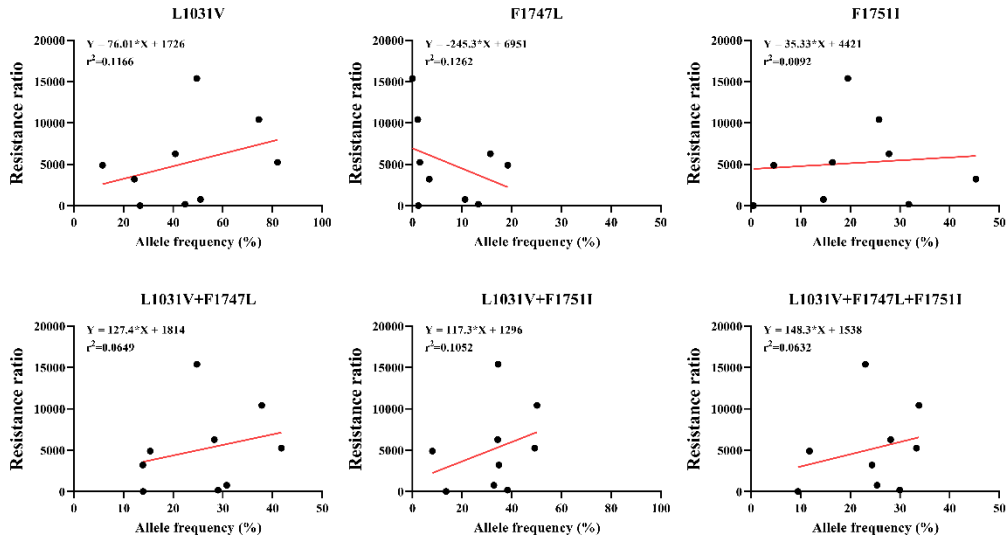


Fig. S3. Bioassays of permethrin resistance in transgenic flies. Data are means \pm S.E.M.

Statistically significant differences are indicated as: *: $p < 0.05$; **: $p < 0.01$; and ***: $p < 0.001$.

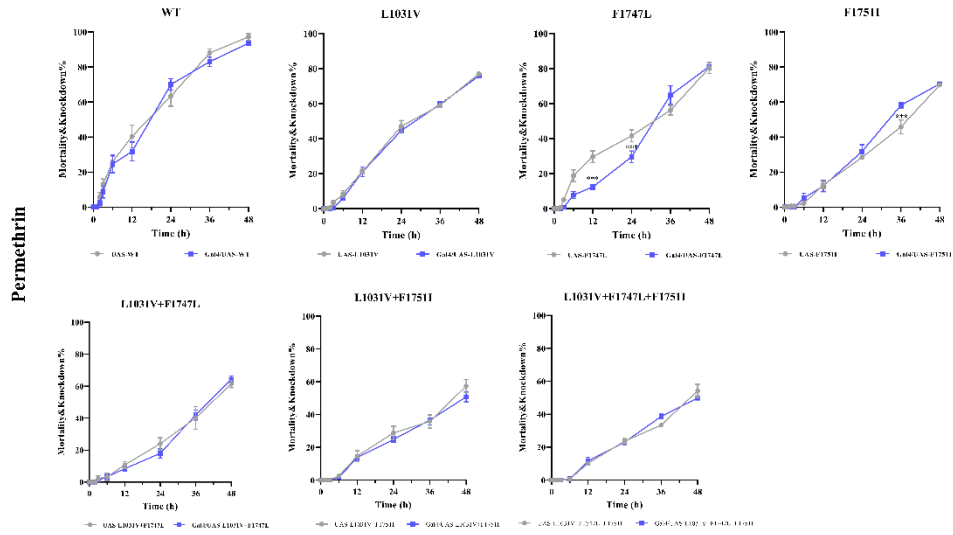


Fig. S4. Bioassays of beta-cypermethrin resistance in transgenic flies. Data are means \pm S.E.M. Statistically significant differences are indicated as: *: $p < 0.05$; **: $p < 0.01$; and ***: $p < 0.001$.

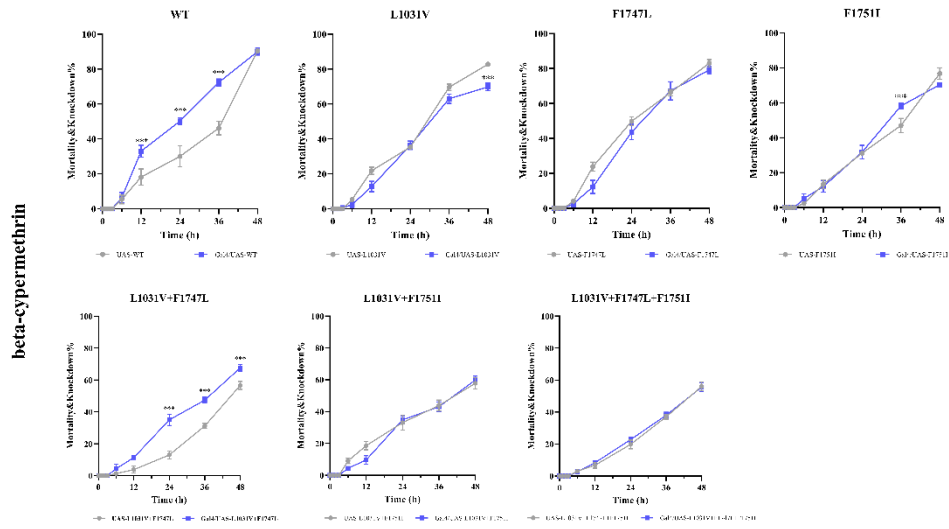


Fig. S5. Pyrethroid docking predictions. Predictions for (A) permethrin, (B) fenpropathrin, and (C) beta-cypermethrin are shown as yellow sticks and residues within 6 Å of docked ligands are shown as grey sticks or are colored green (residue 1747), orange (1751) or brown (1031). Bottom panels (D-F) show the same docking prediction for each pyrethroid as corresponding top panels, but with the channel rendered in surface view.

