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2 **Running title: Pyrethroid resistance of brown dog ticks**

3 **Permethrin Resistance in Field Populations of *Rhipicephalus Sanguineus* Sensu Lato**
4 **(Latrielle, 1806) Collected from Dogs in eastern of Iran**

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20 **ABSTRACT**

21 The high level of acaricide resistance in ticks becomes a challenge for dog owners in Iran.
22 This study was conducted in South Khorasan province of Iran at 2024. In this study, the
23 resistance status of *Rhipicephalus Sanguineus* (Acari: Ixodidae) to permethrin at various
24 concentrations were evaluated using the Larval Packet Test (LPT) method recommended by
25 the Food and Agriculture Organization (FAO). PCR assays were conducted to investigate the
26 mechanisms of resistance to acaricides. We used PCR to amplify segment 6 of domain III of
27 the voltage-sensitive sodium channel protein from both pyrethroid-susceptible and pyrethroid-
28 resistant tick strains. The LPT discriminating dose bioassays confirmed the pyrethroid
29 resistance phenotype of the analyzed strains. The mortality rate at LC₉₉ was ranged between
30 38.1 to 49.1%. At discriminating dose, survival rates ranged from 48.3% to over 60.1%.
31 Additionally, of the 40 ticks analyzed, mutations C2130T and T2134C were detected in 38
32 (95%) ticks. The presence of permethrin resistance in *R. sanguineus* s.l. populations in Iran
33 highlights the need for alternative control strategies, and the identification of genetic mutations
34 provides valuable information for understanding the mechanisms of resistance.

35 **Keywords:** *Rhipicephalus sanguineus*; acaricide resistance; diagnostic concentration;
36 permethrin

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40 1.INTRODUCTION

41 Ticks are one of the most important arthropod vectors of disease-causing agents in both humans
42 and animals. The *R. sanguineus* is an important tick species that feeds mainly on dogs but can
43 also infested other mammalian hosts. ⁽¹⁾ *R. sanguineus* feed on the blood of their hosts and
44 transmit a wide range of pathogens, including viruses, bacteria, and protozoans. ⁽²⁾ *R.*
45 *sanguineus*, the most commonly found tick around the world due to its biological flexibility.
46 One of the primary methods of controlling tick infestations is through the use of acaricides.
47 However, the excessive and often inappropriate use of acaricides has led to the emergence of
48 acaricide resistance, including *R. sanguineus*. ^(3, 4) Understanding the probable acaricide
49 resistance in *R. sanguineus* populations in Iran is crucial for developing effective strategies to
50 control tick infestations and prevent the transmission of tick-borne diseases. ⁽⁵⁾ Acaricide
51 resistance is a complex phenomenon that involves various genetic and physiological
52 mechanisms. These mechanisms can result in decreased sensitivity to the acaricides used to
53 control tick populations. ⁽⁶⁾ Recent studies have suggested that acaricide resistance in tick
54 populations is multifactorial and involves several mechanisms, including target-site
55 insensitivity, metabolic detoxification, and changes in behavior and physiology. ⁽⁷⁾ Target-site
56 insensitivity involves mutations in the genes that code for the target sites of the acaricides,
57 resulting in decreased binding of the acaricides and reduced effectiveness in killing the ticks.
58 Metabolic detoxification involves the overexpression of enzymes that can break down the
59 acaricides, making them less effective. Changes in behavior and physiology involve alterations
60 in the tick's behavior, such as reduced exposure to the acaricides, and changes in the tick's
61 physiology, such as altered cuticle permeability, which can reduce the uptake of the acaricides.
62 The emergence of acaricide resistance in *Rhipicephalus* populations in Iran is a major concern
63 for both animal and public health(8). Further research is needed to elucidate the molecular and
64 physiological mechanisms underlying acaricide resistance in *R. sanguineus* populations in Iran.

65 2. MATERIAL AND METHODS

66 2.1. Sample Collection

67 During June 2022 to May 2023, brown dog ticks were collected from sheepdog of four
68 locations in rural areas located in South Khorasan provinces, east of Iran. The engorged and/or
69 partially engorged female ticks were collected from naturally infested dogs using tick
70 infestation methods, tick drags, and visual searches. The collected ticks were transported
71 immediately to the laboratory in vials containing moist filter paper. The morphological
72 identification of collected samples were confirmed under a stereo-microscope using the
73 standard keys ⁽⁹⁾. From each colony, 30 engorged females were incubated in an environmental
74 chamber at 26–27 °C and 85±5% relative humidity for 3-4 weeks to allow egg lying. The 14-
75 21 day old tick larvae were utilized for the bioassay experiments. The female adult specimens
76 that had been depleted of eggs were isolated, rinsed with distilled water, and then dried using
77 paper towels. Each individual was then frozen separately at a temperature of -80°C for future
78 use in molecular analysis.

79 2.2 Acaricide bioassays

80 The sample size calculation was based on WHO guideline (10).The efficacy of permethrin was
81 assessed using the larval packet test (LPT) developed for acaricide testing of tick populations.
82 ⁽¹¹⁾ Technical-grade 92% permethrin (Mumbai, India) were used as the active ingredients for
83 the LPT. A stock solution was prepared by dissolving permethrin in a 2:1 ratio using
84 trichloroethylene (TCE) (Merck, Germany), and olive oil. ⁽¹²⁾ In Iran, the standard susceptible
85 indigenous strain of *R. sanguineus* was not available. Therefore, in this study, the
86 discriminating concentration of acaricide-susceptible brown dog tick strain was acquired from
87 previous study that was set as 0.19% . ⁽¹³⁾The DC used was calculated by doubling the lethal
88 concentration 99.9% (LC99) derived from a series of tests conducted with a susceptible strain.

89 ⁽¹⁴⁾The LC99 of 0.09% active ingredient (AI) was also tested. Bioassays were conducted on
90 three replicates with 100 larvae per pocket for each concentration.

91

92 2.3 Molecular analysis

93 The genomic DNA of 10 *R. sanguineus* larvae from each location was extracted using the
94 DNeasy® Blood and Tissue Kit (QIAGEN) as the manufacturer's guidelines. Each larva was
95 homogenized in 50 microliters of distilled water and incubated at 56°C for 6 hours before being
96 transferred to the column for preparation. The quality and concentration of the DNA obtained
97 were assessed through agarose gel electrophoresis and a Nanodrop spectrophotometer. PCR
98 amplification was conducted in a total volume of 25 µl, containing 2 µl of template DNA, 1 µl
99 of each primer (forward and reverse primers), 12.5 µl of 2X Taq PCR MasterMix (Takara,
100 Japan), and 8.5 µl of nuclease-free water. The primers FG-228 (5'- CTA ACA TCT ACA TGT
101 ACC -3') and BDT-227 (5'- TTG TTC ATT GAA ATT GTC AA-3') were utilized for
102 amplification of the domain III segment VI of the sodium channel gene. ⁽¹⁵⁾ The PCR
103 amplification was carried out with an initial denaturation at 96°C for 3 min, followed by 35
104 cycles of denaturation at 94°C for 1 min, annealing at 50°C for 1 min, extension at 72°C for 1
105 min, and a final extension at 72°C for 7 min. In total, 20 samples demonstrating phenotypic
106 susceptibility and 20 samples displaying phenotypic resistance were used for the sequence
107 analysis.

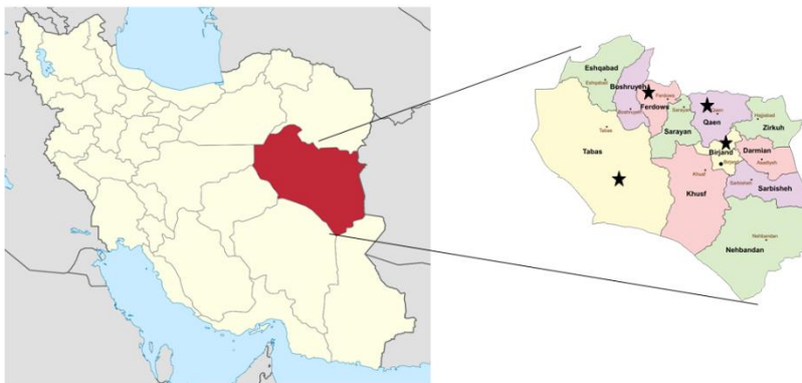
108 2.4 Statistical Analysis.

109 The evaluation of mortality was conducted at 24 hours. The adjust of control mortality was
110 calculated based on the formula of Abbott. ⁽¹⁶⁾ The percentage survival was recorded for each
111 multiple of the diagnostic concentration. The classification of resistant phenotypes will be

112 placed in three classes: low resistance (60 to 90% mortality in LC99×2), moderate resistance
113 (13 to 50% mortality in LC99×2), and severe resistance (1 to 12 Mortality percentage in
114 LC99×2).⁽¹⁷⁾

115 3. RESULTS

116 This study represents the initial assessment of acaricides efficacy on *R. sanguineus* population
117 in South Khorasan provinces (Figure 1).



118
119 FIGURE 1. The collection site of ticks were shown.

120
121 Of these study, Only 4 population of *R. sanguineus* were reared successfully and provided
122 sufficient numbers of larvae and subsequently subjected to bioassay to test their susceptibility
123 to permethrin. The field cached *R. sanguineus* strains were evaluated for mortality with
124 permethrin concentrations 1 and 2 times the diagnostic concentrations, i.e. 0.09 and 0.19%. The
125 mortality rate at LC99 was ranged between 40.5 to 49.1% (Table.1).

126 Table 1. The average lethal rate of *Rhipicephalus sanguineus* (Latreille) strains, collected from
127 various regions in the east of Iran, when exposed to permethrin

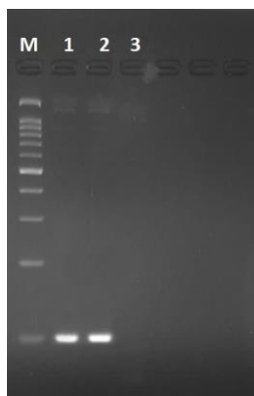
128

Strain	Location	LC ₉₉ (0.09% AI) Mortality	2×LC ₉₉ (0.19% AI) Mortality
B1	Birjand	40.5	49.6
B2	Ferdows	42.5	48.3
B3	Ghaen	49.1	60.1
B4	Tabas	38.1	65.1

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131 At 2×LC₉₉ (0.19% AI), lethal rates ranged from 48.3% to over 65.1%. To screen for mutations
 132 on the sodium channel's domain III segment VI, sequencing was conducted on 10 random
 133 samples from each phenotypically resistant population of brown dog ticks (Figure 2).



134

135 FIGURE.2 Agarose gel separation of representative PCR products of the voltage-sensitive sodium
 136 channel gene. Lane 1–2, positive isolates; Lane 3 negative control, DNA ladder 100 bp

137

138 The analysis revealed four genotypes on domain III among the *R. sanguineus* population from
 139 east of Iran by comparing the susceptible (GenBank KU886031) and permethrin-resistant

151 FIGURE 4. Gene sequences of the voltage-gated sodium channel of *Rhipicephalus sanguineus*
152 aligned with that of wild sequences (GenBank accession number:KU886031) showing the
153 mutations C2130T and T2134C .Three haplotypes were reported.

154

155 **4. DISCUSSION**

156 This study provides the first laboratory-confirmed permethrin resistance data for brown dog ticks from
157 the East of Iran. *R. sanguineus* is one of the the most prevalent infected ticks for different
158 pathogen in Iran. ⁽¹⁸⁾ The results of this study provide important preliminary insights into the
159 efficacy of permethrin on the *R. sanguineus* population in east of Iran. The findings show that
160 the mortality rates of *R. sanguineus* populations varied significantly when subjected to different
161 concentrations of permethrin. At 2×LC₉₉ (0.19% AI), lethal rates ranged from 48.3% to over
162 65.1%, indicating that this concentration is not effective for controlling of the following tick
163 population. Previous studies in Iran have also shown high levels of resistance to pyrethroid
164 insecticides among populations of *Rhipicephalus* ^(8, 19). Limited studies have been carried out
165 on the resistance of ticks to pyrethroid in Iran, ^(20, 21) and the present study is the first
166 comprehensive investigation of the *R. sanguineus* in this area. Previous studies from around
167 the world also showed resistance to pyrethroid pesticides among *R. sanguineus*^(12, 13).
168 Importantly, our bioassay findings highlight the need for careful consideration of appropriate
169 concentrations of acaricides to achieve effective tick control, and suggest that higher
170 concentrations may be necessary to achieve satisfactory results. Overall, these results constitute
171 an important step towards the development of more effective and targeted approaches for tick
172 control in Iran.

173 Of these study, Only 4 population of *R. sanguineus* were reared successfully. An important
174 consequence of resistance development in tick populations may be a decline in overall fitness.

175 According to Roma et al. (2010), exposure to sub-lethal levels of permethrin adversely affects
176 reproductive success (22). Subsequent research could explore how these sub-lethal
177 concentrations of permethrin impact the reproductive capacity of adult female *R. sanguineus*
178 with SNPs in comparison to their susceptible counterparts.

179 The current study identified a mutation on domain III segment VI of the sodium channel that
180 was responsible for resistance to insecticides in the tick population. ^(3, 23) In previous studies, it
181 has been shown that T2134C mutations in this gene is associated with resistance to pyrethroid
182 resistance in *R. sanguineus*. ⁽³⁾ The findings reveal that out of the 40 ticks examined, just 5%
183 were wild strains, suggesting that the majority of ticks had been subjected to selection pressure
184 and had acquired resistance to insecticides. In this study, 38 out of 40 samples (90%) carried
185 the T2134C mutation that could be the explained the high levels of permethrin resistance.
186 However, it is possible that other mechanisms, such as metabolic detoxification, sequestration,
187 reduced penetration, or additional mutations in the sodium channel, may be related to
188 insecticide resistance. ^(24, 25) Overall, this study underscores the importance of bioassay and
189 genetic studies in understanding and controlling brown dog ticks populations. The number of
190 samples collected may not fully represent the genetic diversity of the tick populations across
191 the eastern regions of Iran. A larger sample size from various geographical locations could
192 provide a more comprehensive understanding of resistance patterns. The study primarily
193 focused on permethrin resistance, which may not reflect the overall resistance profile of the
194 tick populations to other classes of acaricides. A broader assessment of resistance to multiple
195 insecticides would provide a more complete picture. Limited funding restricted the scope of
196 the sequencing project, potentially leading to a smaller sample size and fewer gene targets
197 being analyzed than initially desired.

198

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203 **Conflict of interest**

204 The author declare no conflict of interest

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206 **Authors' Contribution:**

207 A. V: Writing – review & editing, Writing – original draft, Project administration,
208 Methodology, Formal analysis, Data curation, Conceptualization. R. S: Writing – review &
209 editing, Writing – original draft, Visualization, Validation, Supervision, Resources,
210 Methodology, Investigation, Formal analysis, Data curation, Conceptualization. E Kh: Writing
211 – review & editing, Writing – original draft, Software, Methodology, Investigation, Formal
212 analysis. S. Sh: Writing – review & editing, Visualization, Validation, Supervision, Resources,
213 Funding acquisition, Conceptualization.

214 **Ethics**

215 Research ethics committee of islamic azad university, science and research branch
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220 **Data Availability**

221 Should there be a need for data that support the findings of this study, they are available from
222 the corresponding author upon reasonable request.

223

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