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High-resolution melt (HRM) analysis for detection of SNPs associated with pyrethroid resistance in the southern cattle fever tick, *Rhipicephalus (Boophilus) microplus* (Acari: Ixodidae)∗

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ABSTRACT

The southern cattle fever tick, *Rhipicephalus (Boophilus) microplus*, is the most economically important ectoparasite of cattle worldwide. A limitation for sustainable control and eradication is the emergence of acaricide resistance among tick populations. Molecular diagnostic tools offer the opportunity to detect resistance rapidly, which can be complemented with confirmatory bioassays with larvae and adult ticks that are more resource and time consuming to generate. Synthetic pyrethroid resistance is one of the most prevalent and well-studied forms of resistance in arthropods, being linked with target site alterations in the sodium ion channel gene. Here, we report research on a novel molecular method to detect mutations in the para-sodium channel gene of *R. microplus* associated with acaricide resistance that is based on quantitative PCR high-resolution melt (HRM) analysis. Genomic DNA fragments of domains II and III of the para-sodium channel gene were amplified by real-time PCR in the presence of Eva™Green dye to test resistant and susceptible reference ticks from the U.S., Brazil, and Mexico. Larval packet tests with discriminating doses and a modified lethal time analysis were performed to confirm resistance to permethrin, cypermethrin, deltamethrin, and flumethrin in laboratory strains. Tick specimens collected from cattle that were inspected at the United States Port-of-Entry at the Texas-Mexico border were also genotyped. Previously described mutations associated with pyrethroid resistance (T170C, C190A, G184C, and T2134A) were successfully detected by qPCR-HRM in different genotypes and confirmed by sequencing. A novel non-synonymous SNP located at domain III (C2136A) and the G215T mutation in domain II, previously described only in Asian *R. microplus* and *R. australis* were also detected with the HRM and confirmed by sequencing. This technique could be adapted for high-throughput screening, detection, and discovery of allele-specific mutations in cattle tick outbreak populations to inform eradication strategies in the USA. This knowledge could also be applied to integrated control programs in other parts of the world where *R. microplus* is endemic and where similar SNPs have been identified associated with pyrethroid resistance. This study highlights the existence of several mutations in the para-sodium channel gene in different combinations in field populations of *R. microplus* from Mexico.

1. Introduction

Ticks infesting cattle and other farm animals in tropical and subtropical areas of the world are controlled primarily through the use of products containing synthetic acaricides. Resistance to one or more classes of acaricides, including pyrethroids, has been reported among...
populations of the southern cattle fever tick, *Rhipicephalus (Boophilus) microplus* (Canestrini), which is considered the most economically important ectoparasite of livestock in parts of the world where it is endemic (Grisi et al., 2014; Rodriguez Vivas et al., 2017). *R. microplus* is also of veterinary relevance because it is a vector of the pathogens that cause bovine babesiosis, anaplasmosis and spirochaetosis (Aubry and Geale, 2011; Pérez de León et al., 2014; Walker et al., 2003).

The characterization of pyrethroid resistance in *R. microplus* and other arthropod pests has been extensively explored (Rinkevich et al., 2012; Lovis et al., 2012; Nogueira Domingues et al., 2012; Kumar et al., 2013; Robbertse et al., 2016; Wyk et al., 2016; Bandara et al., 2009; Aguirre et al., 2010; Jonsson et al., 2010; Rodriguez-Vivas et al., 2012; Lovis et al., 2012; Nogueira Domingueas et al., 2012; Kumar et al., 2013; Robbertse et al., 2016; Wyk et al., 2016; Bandara and Karunarante, 2017; Sungirai et al., 2018). In addition to target site insensitivity, other mechanisms involved in pyrethroid resistance among *R. microplus* populations include detoxification enzymes such as esterases (Guerrero et al., 2002; Singh and Rath, 2014; Gupta et al., 2016) and monooxygenases (Graham et al., 2016). Bioassays like the larval packet test (Stone and Haydock, 1962), or the adult immersion test (Drummond et al., 1973) are used extensively to detect acaricide resistance phenotypes. These functional bioassays can discriminate the levels of resistance between tick populations and can be adapted to investigate possible mechanisms of resistance (Li et al., 2003). However, these bioassays require access to abundant numbers of viable engorged females that need to be collected, maintained for lengthy periods of oviposition and testing of viable larvae. Often, this can be difficult to achieve especially in areas of low infection levels or where strict tick control measures are practiced.

Molecular methods to detect acaricide resistance can be performed with few ticks, including samples of properly conserved dead ticks (e.g. in ethanol or isopropanol). The first description of an allele-specific polymerase chain reaction (AS-PCR) assay to genotype pyrethroid-resistant tick strains was reported by Guerrero et al. (2001). This AS-PCR assay detects a mutation in domain III (T2134A) and was applied to studies on the epidemiology of pyrethroid resistance in *R. microplus* from Mexico (Guerrero et al., 2002; Rosario-Cruz et al., 2009; Rodriguez-Vivas et al., 2012). Lovis et al. (2012) proposed a multiplex AS-PCR, aimed to detect three (C190A; G215T and T2134A) single nucleotide polymorphisms (SNPs) in the *para*-sodium channel gene. More refined techniques were also developed, as a ‘Tagman®’ dual probe quantitative PCR diagnostic assay to detect the C190A mutation (Morgan et al., 2009), and a melt analysis of mismatch amplification mutation assay (melt-MAMA) qPCR platform, designed to detect the mutations T170C, C190A, and T2134A (Stone et al., 2014).

High-resolution melt analysis (HRM) is a closed-tube post-PCR application involving the use of a dye that fluoresces when intercalated with double-stranded DNA (dsDNA), but not when in solution after the denaturation of DNA (Reed et al., 2007). The intercalated dye is released when dsDNA denatures, resulting in a loss of fluorescence at the melting temperature of the PCR product. HRM curves are generated by measuring the decrease of fluorescence as the temperature is slowly increased with a high level of accuracy (0.05–0.1°C). The fluorescence decreases at each step due to the transition of the DNA from double- to single-stranded. The melting temperature of a DNA molecule is determined by its nucleic acid sequence and length (Reed et al., 2007). Differences in these nucleotide sequences between samples result in melting profiles that are unique to particular genotypes, allowing for differentiation. A PCR reaction that contains only one type of DNA sequence, such as homozygous templates from diploid organisms, produces a melt curve with one peak. Heterozygous templates result in a mixture of homopolymers and heteropolymers. Due to the imperfect binding of their strands, the melting temperature is strongly lowered in the heteropolymers, resulting in an early peak in the curve. Multi-loci SNP products will cause more complex heteropolymer formations and additional peaks in the melt curves (Mader et al., 2008).

HRM was successfully used to detect pyrethroid resistance mutations in the yellow fever mosquito *Aedes aegypti* (Wulianjadi et al., 2015), and in the scabies mite *Sarcoptes scabiei* (Passay et al., 2008). In a direct comparison with other techniques (i.e. AS-PCR and Taqman probes) the HRM had excellent performance in detecting knock down resistance (*kdr*) mutations in previously sequenced *Anopheles gambiae* standard samples. However, when the technique was used to detect those mutations in field derived samples with variable quantity and quality of DNA, its performance was not as efficient as the Taqman probes. Nevertheless, the cost of the analysis per sample was lower than the other two techniques (Bass et al., 2007).

Here, we developed an HRM assay to simultaneously detect multiple mutations in the *para*-sodium channel gene associated with pyrethroid-resistant *Rhipicephalus microplus*. Reference laboratory tick strains from Mexico, Brazil and USA, as well as field samples from Mexico were used in this study.

### 2. Materials and methods

#### 2.1. Ticks

Colonies of *R. microplus* maintained at the USDA-ARS Cattle Fever Research Laboratory (CFTRL) in Edinburg, Texas were used in this study and were reared as described by Davey et al. (1980). The Deutch strain served as the acaricide susceptible reference collected originally in Laredo, Texas and maintained as a colony at the CFTRL since 2001. Deutch strain ticks from the F29 generation were used in this study. Santa Luiza is the colony that includes ticks resistant to permethrin and amitraz, which were collected from a ranch in Rio Grande do Sul, Brazil; this colony was maintained at the Mexican National Parasitology Laboratory, Juípete, Morelos, Mexico, before being established at the CFTRL in 2000 (Li et al., 2004). Ticks from the F64 generation of the Santa Luiza colony were used. Ticks from the El Zamora colony are resistant to permethrin, amitraz, and fipronil, and were collected in 2010 from a ranch in the State of Tamaulipas, Mexico (Miller et al., 2013). F59 generation of El Zamora ticks were used. Ticks to establish the Yucatan colony were collected in 2014 from red deer (*Cervus elaphus*) at a ranch located in the State of Yucatan, Mexico (Rodriguez-Vivas et al., 2014), and are resistant to cypermethrin, coumaphos, and ivermectin. Ticks of the El Zamora F41 generation were used.

*Rhipicephalus microplus* field samples were obtained from cattle during inspections at the United States Ports-of-Entry located at the border of Mexico and the State of Texas. Ticks were received at the CFTRL inside 5 mL plastic assay tubes and were stored in cryovials and immediately frozen at −80°C to be used in the molecular analysis. In total, 29 semi-engorged females were processed and those were taken from cattle with origins in different states of Mexico: Coahuila (*n* = 3); Nuevo León (*n* = 7); Tamaulipas (*n* = 14) and Veracruz (*n* = 5). No previous information about the pyrethroid resistance status was obtained from those ticks.

#### 2.2. Preparation of ticks

Engorged female *R. microplus* from the susceptible (Deutch), and resistant colonies (Santa Luiza, El Zamora, and Yucatan) were collected after their natural detachment from cattle according to the protocol approved by the Institutional Animal Care and Use Committee. After collection, the ticks were washed with water and dried with paper towels. From each colony, 30 engorged females were incubated in an environmental chamber at 28°C, and a relative humidity of 92% in a plastic petri dish (9 cm diameter) for 20 days to allow egg laying (Davey et al., 1980). Depleted females were separated from the eggs, washed with distilled water, dried with paper towels and individually frozen at
− 80 °C to be used in the molecular analysis. The eggs were mixed thoroughly and incubated under the same conditions in 2-dram glass vials closed with cotton plugs to allow the passage of air and moisture to permit larval hatching. The egg masses were checked daily for larval hatching. Larvae 14–21 days old were used in the bioassays.

2.3. Bioassays

The bioassays were performed with analytical standards of the following chemicals: permethrin (FMC, Philadelphia, PA, USA), cypermethrin, deltamethrin, and flumethrin (Sigma Aldrich Co., St. Louis, MO, USA). In order to characterize the phenotypic resistance to these chemicals, the larval packet test procedure was used with pre-established discriminating doses (DDs) calculated as $2 \times \text{LC}_{50}$ of a susceptible reference strain. The DDs were 0.25% permethrin (Miller et al., 1999), 0.20% cypermethrin, 0.06% deltamethrin, and 0.01% flumethrin (FAO, 2004). Acaricides were diluted in a mixture containing two parts of trichloroethylene and one-part olive oil (TChE-OO) in order to prepare the impregnation solutions. A volume of 0.67 mL of each acaricide solution was used to impregnate a piece of quantitative filter paper (85 mm × 75 mm) - Omnifit, Mal-dstone, England). The material was left to dry for 2 h inside a fume hood to allow for trichloroethylene evaporation. After drying, packets of the same acaricide were wrapped in aluminum foil and maintained at 4 °C until used.

On the day of testing, filter papers were taken from the refrigerator, folded in the middle, and sealed on both sides with metal clips to form the packets. Approximately 100 tick larvae were transferred to each packet using a flat paintbrush. The packets were sealed with a third clip on top, and incubated at 27 ± 1 °C and 80–90% relative humidity. Control groups were exposed to filter papers impregnated with acaricide-free TChE-OO. After 24 h, larval mortality was determined by counting the total number of dead and viable individuals. Larvae that were paralyzed or moving only their appendices without the ability to walk were considered non-viable. Three packets impregnated with each acaricide as well as controls were prepared for each tick sample, i.e. from the Deutch, Santa Luiza, El Zamora, and Yucatan colonies.

Percentage of larval mortality was determined for each of the three larvae that were incubated inside a fume hood to allow for trichloroethylene evaporation. After drying, packets of the same acaricide were wrapped in aluminum foil and maintained at 4 °C until used.

On the day of testing, filter papers were taken from the refrigerator, folded in the middle, and sealed on both sides with metal clips to form the packets. Approximately 100 tick larvae were transferred to each packet using a flat paintbrush. The packets were sealed with a third clip on top, and incubated at 27 ± 1 °C and 80–90% relative humidity. Control groups were exposed to filter papers impregnated with acaricide-free TChE-OO. After 24 h, larval mortality was determined by counting the total number of dead and viable individuals. Larvae that were paralyzed or moving only their appendices without the ability to walk were considered non-viable. Three packets impregnated with each acaricide as well as controls were prepared for each tick sample, i.e. from the Deutch, Santa Luiza, El Zamora, and Yucatan colonies.

Percentage of larval mortality was determined for each of the three packets of each acaricide treatment. For each test, mean mortality and standard errors were calculated with Microsoft Excel (Microsoft Corporation, Redmond, WA).

The knockdown effect of permethrin was assessed with a lethal time bioassay. About 100–150 larvae of each strain were incubated inside the filter paper packets impregnated with 0.25% permethrin as described above. The mortality was determined at six time points: 10, 20, 30, 40, 50, and 60 min after the exposure, as described for the resistance test above. Mortality data were analyzed using the Probit model in the Polo Plus software (Version 1.0, licensed. LeOra Software, 2003). For each test, the following parameters were determined: median lethal time (LT50), with its respective 95% confidence limits (95% CL), and the slope of the regression line. The resistance ratios (RR) were obtained using the Polo-Plus software employing the formula described by Robertson et al. (2007). Comparisons were determined to be significant when the calculated 95% CL did not overlap. For visualization of data, the log time vs. probit converted mortality plots were generated using Microsoft Excel (Microsoft Corporation, Redmond, WA).

2.4. DNA extraction

The genomic DNA of ticks was extracted using a phenol-chloroform method. Frozen ticks were taken from ultracold freezer (− 80 °C) and transferred to 2-mL plastic tubes containing five ceramic beads (2.8 mm diameter - Omni International, Kennesaw, GA, USA) each. The tubes also contained 600 μL of lysis buffer (10 mM Tris-HCl pH 8; 2 mM EDTA pH 8; 0.5% SDS), 3 μL proteinase K (20 mg/mL, Invitrogen, Carlsbad, CA, USA), and were processed at 4000 rpm for 15 s with a bead mill homogenizer (Omnim International, Kennesaw, GA, USA). Homogenates were incubated for 12–18 h at 55 °C to allow for protein digestion. Following, the samples were incubated at 65 °C for 15 min to inactivate the proteinase K. After briefly cooling in an ice bath, 5 μL of RNase A (20 mg/mL, Invitrogen, Carlsbad, CA, USA) was added to the homogenates followed by incubation at 37 °C for 15 min to remove RNA contamination. The homogenates were submitted to a phenol-chloroform DNA extraction procedure. The DNA was precipitated in an ice-cold absolute ethanol and sodium acetate solution at − 20 °C for 12–18 h. The DNA pellets were collected by centrifugation (10,000 g at 4 °C for 15 min) and washed two times with 70% ethanol. The final pellet was suspended in 50 μL Tris-EDTA, pH 8. The genomic DNA was quantified on an EON spectrophotometer (Biotek, Winooski, VT, USA), and diluted to 100 ng/μL for PCR.

2.5. PCR and sequencing

In order to identify the mutations, we used primers designed to amplify the exon region of the para-sodium channel gene domain II S4−S5 linker and the exon region of the domain III S6. The identification of the SNPs present in domains II and III was obtained by Sanger dideoxy sequencing.

Sequences of the exon encoding domain II were obtained from DNA fragments amplified with the primers designed by Morgan et al. (2009) where the forward BmNaF5 (5'TACGTGTTGTTCAAGCTAGC) and reverse primer BmNaR5 (5'ACTTCTTCTGTAGTCTTGC) yield a 167 bp product. PCRs were carried out in 50 μL volumes containing the following reagents: 100 ng of DNA template, 1 × PCR buffer, 1.5 mM MgCl2, 0.2 mM dNTPs, 200 nM each primer and 1 U of AmpliTaq Gold II polymerase (Applied Biosystems, Carlsbad, CA, USA). The reaction was performed under the following conditions: 95 °C for 5 min, followed by 40 cycles of 95 °C for 30 s, 50 °C for 30 s, 72 °C for 30 s, and a final extension step of 72 °C for 7 min and final hold at 4 °C.

Sequences of the exon encoding domain III were obtained from DNA fragments amplified with the primers described by Stone et al. (2014). Initially a 135 bp fragment was amplified using the forward primer RmNaDomainIIIF1 (5'AAGAGGACAAACCGGATAGC) and reverse primer RmNaDomainIIIR2 CON (5'CTTCTTTGTGTTAGTGAAT TGT). PCRs were carried out in 10 μL volumes containing 100 ng of DNA template, 1 × PCR buffer, 2.5 mM MgCl2, 0.2 mM dNTPs, 1 U of AmpliTaq Gold II polymerase. The reaction was performed under the following conditions: 95 °C for 10 min, followed by 40 cycles of 94 °C for 60 s, 53 °C for 30 s, 72 °C for 30 s. The PCR products from this first reaction were diluted 1:1000 and used as the template for a second PCR using tailed primers. The second PCR used forward tailer primer RmNaDomainIIIF3 (5'acccacatgagagcAAGAGGACCGGAAT TCG) and reverse tailed primer RmNaDomainIIIR3 (5'acgacct gactgcttcTTCTTTTGTTTCAGTGAAT TGT) resulting in an amplicon length of 173 bp. The conditions of the second PCR were identical to the first, with exception of the annealing temperature that was 65 °C.

PCR products were visualized by electrophoresis on 2.2% agarose FlashGel DNA casettes with a FlashGel System (Lonza Rockland, Inc., Rockland, ME, USA). PCR products were purified using Diffinity RapidTip (Chiral Technologies, Inc., West Chester, PA, USA), and sequenced using the primers BmNaF5 and BmNaR5 for the domain II amplicons and tail primers RmNaDomainIIIF3seq (5'acccacactag agacgacAGAGGACCGGAAT TCG) and RmNaDomainIIIR3seq (5'acgacctgactgcttc) for amplification through capillary electrophoresis in a 96-capillary Applied Biosystems 3730xl DNA Analyzer (Retrogen, Inc.).

The validation of the allele sequences for heterozygous individuals was performed by cloning and sequencing of PCR amplicons. PCR products were ligated into the pJET 1.2/blunt vector of the CloneJET PCR Cloning Kit (Thermo Fisher Scientific, Waltham, MA, USA) according to the manufacturer’s specifications. Ligations were used to transform DH5α competent cells (Invitrogen, Carlsbad, CA, USA). Up to
eight colonies per individual tick were selected for PCR screening with the pJET1.2 plasmid primers. Six clones containing the correct insert size were chosen for sequencing according to the methods described above using pJET1.2 primers. Sequencing data were aligned and translated with CLC Main Workbench 7 (Qiagen Aarhus A/S, Copenhagen, Denmark). Sequences were compared to each other and analyzed for the presence of the different SNPs.

2.6. Detection of domain III mutation

Previously described AS-PCR technique was used to genotype the ticks for the T2134A mutation (Guerrero et al., 2001, 2002). PCR was performed using 20 µL reactions containing the following reagents: 100 ng of DNA template, 1 × PCR buffer, 1.75 mM MgCl2, 0.2 mM dNTPs, 100 nM each primer and 0.5 U AmpliTaq Gold II polymerase (Applied Biosystems, Carlsbad, CA, USA). The reactions were carried out in a thermocycler programmed for 96 °C for 2 min followed by 37 cycles, each consisting of denaturation at 94 °C for 1 min, annealing at 60 °C for 1 min, and extension at 72 °C for 1 min. The program also included a final extension step at 72 °C for 7 min. PCR products were visualized by electrophoresis on 2.2% agarose as described above.

2.7. Quantitative PCR - HRM assays

Two high-resolution melt (HRM) assays were developed to genotype the different pyrethroid resistance related polymorphisms in the para-sodium channel gene of R. microplus. qPCR primers were designed using PRIMER3WEB v.4.0.0 (Untergasser et al., 2012). The first assay targeted the domain II S4–S5 mutations, and the primers were designed to flank the following nucleotide substitutions: T170C, G184C, C189A (silent), C190A and G215T. Reactions with the forward primer HRM_Rm_Na-D2_Fw (5′ GATTCCCAGGACAAAGGTCA) and the reverse primer HRM_Rm_Na-D2_Rv (5′ GATTGCCACGCAAGGGTC) yielded an 89 bp amplicon. The second assay targeted the Domain III S5 mutation T2134A. Primers used on the reactions were HRM_Rm_Na-D3_Fw (5′ GATTGCCACGCAAGGGTC) and HRM_Rm_Na-D3_Rv (5′ GATTGCCACGCAAGGGTC) which produced a 61 bp amplicon.

Twenty-four ticks from each colony and the ticks from the field collections were genotyped and the reactions were performed in duplicates. Each run contained the samples, a non-template control (water), clones corresponding to wild-type and mutant genotypes and homozygous and heterozygous genomic DNA references. PCR reactions (10 µL) contained 1X Precision Melt Supermix (Biorad, Hercules, CA, USA), 200 nM each primer, and 100 ng of genomic DNA template. Reference plasmids were used at 1 ng per reaction. Samples were run on a Mic qPCR Cycler (Bio Molecular Systems Pty Ltd., Upper Coomera, Australia) using the temperature cycling conditions of 95 °C for 2 min to activate the hot-start polymerase, 44 cycles of a denaturation step of 95 °C for 10 s, and annealing/extension at 60 °C for 30 s. Fluorescence information was captured at the end of each 60 °C step. PCR products were then subjected to HRM analysis. The HRM step involved heating the PCR products to 95 °C for 30 s, cooling to 60 °C for 1 min, and then increasing the temperature to 65 °C. The temperature increased from 65 to 95 °C at a rate of 0.08 °C/s, and the change in fluorescence of EvaGreen® was recorded continuously on the green channel of the Mic qPCR Cycler.

Melt curves were generated in the “High Resolution Melt” analysis module of miqPCR software (version 2.6.2, Bio Molecular Systems Pty Ltd., Upper Coomera, Australia). The parameter settings for melt curve normalization of the domain II fragment were: Pre-melt slider = 75.8–77.3 °C, and Post-melt slider = 84.5–86 °C. For domain III, the parameter settings were: Pre-melt slider = 66–67.6 °C, and Post-melt slider = 77.4–80.4 °C. The samples were assigned to a given genotype by examining normalized and difference melt plots in comparison to reference genotypes curves at a 95% confidence limit.

### Table 1

<table>
<thead>
<tr>
<th>Strain</th>
<th>Mean percentage of mortality after 24 h (standard deviation)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.25%</td>
</tr>
<tr>
<td></td>
<td>Permethrin</td>
</tr>
<tr>
<td>Deutch</td>
<td>100 (0)</td>
</tr>
<tr>
<td>Santa Luiza</td>
<td>59.15 (3.53)</td>
</tr>
<tr>
<td>El Zamora</td>
<td>28.12 (3.77)</td>
</tr>
<tr>
<td>Yucatan</td>
<td>2.85 (2.35)</td>
</tr>
</tbody>
</table>

3. Results

#### 3.1. Pyrethroid resistance

The larval packet discriminating dose bioassays confirmed the pyrethroid resistance phenotype of the analyzed strains. Table 1 presents the mortality data of each strain against permethrin, cypermethrin, deltamethrin and flumethrin.

![Fig. 1](image1.png) Fig. 1 and Table 2 shows the results of the lethal time bioassay with permethrin carried out with the four strains. The median lethal time (LT50) for Deutch was 21.74 min (95% CL = 18.48–24.63). After 40 min of exposure, approximately 95% of the larvae of the susceptible strain were dead. For the Santa Luiza resistant strain, the calculated LT50 was 65.02 (95% CL = 53.99–86.51), which is significantly higher than for the Deutch strain. It was not possible to calculate the LT50 for the El Zamora and Yucatan strains, as the observed mortality were found to be at very low rates (~5%) across the observed data points.

#### 3.2. HRM genotyping of reference strains

The HRM assay was successfully used to determine the presence of previously described pyrethroid resistant SNPs in two regions of the para-sodium channel gene of R. microplus (domain II S4–S5 and domain III S6). In domain II, three SNPs were found among individuals of the tick colonies and were validated by cloning and sequencing: G184C, C189A, and C190A. The substitutions in nucleotide 189 were found only in the Coorong, Halls Creek, and El Zamora strains, as the observed mortality were found to be at very low rates (~5%) across the observed data points.

![Fig. 1](image2.png) Figs. 3–5 show the normalized and difference melt plots of the fragments amplified by qPCR. On domain II, eight genotypes could be identified among the laboratory strains. All the ticks from the susceptible reference strain Deutch presented the same genotype (wt), with no substitutions of nucleotides in comparison to the reference sequence (GenBank: AF134216). The C190A and G184C substitutions were found to be associated with both homozygous and heterozygous alleles (Fig. 3A–D). The substitution in nucleotide 189 was found only in heterozygosity and always associated with C190A (Fig. 3A). The occurrence of G184C and C190A was also found among the tick samples, only in heterozygosity (Fig. 5A). On domain III, three genotypes were detected among the laboratory strains: wild type, heterozygous
(T2134W), and homozygous (T2134A) (Fig. 4A and B), and the genotypes were confirmed with the AS-PCR technique and sequencing (Fig. 2B).

The designed HRM assay showed versatility in detecting a number of different genotypes. With two reactions (aimed at domains II and III), we were able to detect and assign six different genotypes (Table 3). One genotype was associated with the susceptible wild-type, and five associated with pyrethroid resistance.

Among the individuals of the El Zamora strain, the only detected mutation was the C190A, with 75% of the ticks being homozygous for this mutation and 25% heterozygous; no wild types were detected. Half of the sampled population (n = 12) presented the C189A silent mutation, always combined with the nucleotide substitution at position 190. All the Yucatan ticks were homozygous for the resistant mutations at domains II (G184C) and III (T2134A) simultaneously.

Ticks from the Santa Luiza strain presented the three SNPs and all the possible genotypes associated with resistance. Most of the Santa Luiza ticks (82.6%) carried at least one resistant allele with a low frequency of resistant homozygous ticks (8.7% for G184C/T2134A and C190A). Five ticks (21.7%) presented the three mutant loci simultaneously. To the best of our knowledge this is the first description of the existence of three simultaneous mutations in the para-sodium channel gene of *R. microplus*.

A good phenotype-genotype correlation of the assays was observed. Pearson Product Moment correlation analysis and linear regression analyses were carried out (Sigma Plot 11.0, Systat Software, 2008) by plotting the frequency of ticks carrying any of the mutations in homozygosis (GG/II/FF and RR/LL/II) against the survivorship in the bioassays with all the pyrethroids tested (Fig. 6). In a Pearson Product Moment correlation analysis, with the exception of cypermethrin ($r = 0.945; p = 0.0552$), there was a significant correlation between resistant-homozygous ticks (RR) and the survivorship after exposure to permethrin ($r = 0.964; p = 0.0361$), deltamethrin ($r = 0.964; p = 0.0363$), and flumethrin ($r = 0.969; p = 0.0311$). This observation was coherent with the results from the bioassays, showing a broad spectrum of resistance against the different pyrethroid acaricides tested (Tables 1 and 2). The high frequency of homozygous resistant ticks in the El Zamora and Yucatan strains was compatible with the "knock-down-resistance" phenotype observed in the lethal-time bioassays (Table 2, Fig. 1). This observation confirmed the association of mutations in the para-sodium channel with resistance to synthetic pyrethroids.

3.3. HRM genotyping of field samples

The developed technique was used with DNA from *R. microplus* field samples from northern states of Mexico (Supplementary table). The HRM was successful in detecting the previously described mutations associated with pyrethroid resistance in the tick colonies (e.g. G184C; C190A and T2134A). Three ticks were wild type for all the loci and can be considered susceptible. Two ticks presented one allele with the G215T mutation, combined with either the T170C or the C190A SNPs

![Fig. 1. Mortality of *R. microplus* larvae of Deutch strain (susceptible) compared to that of the Santa Luiza, El Zamora and Yucatan strains (resistant) exposed to 0.25% permethrin impregnated filter papers. Mortality was evaluated at 10-min intervals for 60 min.](image)

**Table 2** Results of lethal time bioassays conducted with susceptible (Deutch) and pyrethroid resistant (Santa Luiza, El Zamora and Yucatan) reference strains of *Rhipicephalus (Boophilus) microplus* against permethrin.

<table>
<thead>
<tr>
<th>Strain</th>
<th>N</th>
<th>Slope (SE)</th>
<th>Chi-square (DF)</th>
<th>H</th>
<th>LT50 (95%CL)</th>
<th>RR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deutch</td>
<td>1467</td>
<td>4.89 (0.23)</td>
<td>112.89 (16)</td>
<td>7.06</td>
<td>21.74 (18.48–24.63)</td>
<td>–</td>
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<td>Santa Luiza</td>
<td>1107</td>
<td>1.75 (0.18)</td>
<td>23.56 (16)</td>
<td>1.47</td>
<td>65.02 (53.99–86.51)</td>
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</tr>
<tr>
<td>El Zamora</td>
<td>1539</td>
<td>0.39 (0.24)</td>
<td>24.31 (16)</td>
<td>1.62</td>
<td>&gt; 60</td>
<td>a</td>
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<tr>
<td>Yucatan</td>
<td>1531</td>
<td>0.83 (0.3)</td>
<td>9.11 (16)</td>
<td>0.57</td>
<td>&gt; 60</td>
<td>a</td>
</tr>
</tbody>
</table>

N = number of individuals; SE = standard error; DF = degrees of freedom; H = heterogeneity on the Chi-square goodness of fit test; LT50 = median lethal time in minutes; 95%CL = 95% confidence limits, RR = resistance ratio. a: not calculated.
Fig. 2. Nucleotide sequence alignment of sodium channel domains II (2A) and III (2B) DNA fragments of *R. microplus* reference sequence (GenBank Accession Number AF134216) with its respective translated amino acid sequences, and clone sequences obtained from pyrethroid resistant ticks: T170C, G184C, C190A, G184C_C190A, C189A_C190A, C190A_G215T, T170C_G215T, T170C_C189A_C190A, T2134A, T2134A_C2136A, C2130T_T2134A, and C2136A. Identical nucleotides are marked with dots. Numbers above sequences are based on the *R. microplus* para-sodium channel. Sequencing primers positions (Domain II: BmNa5F and BmNa5F; Domain III: RmNaDomainIIIF1 and RmNaDomainIIIRS2-CON) are identified as gray arrows. HRM primers (Domain II: HRM_Rm_Na-D2_F2 and HRM_RmNa_F; Domain III: HRM_Rm_Na F and HRM_RmNa_F) are identified in black. Replaced amino acid residues are identified in red with one asterisk. The synonymous substitutions are identified in orange with two asterisks.
This is the first detection of the G215T, in the Neotropical region. This SNP results in a Gly to a Val change at the amino acid residue 72 in the domain II (Fig. 2A). A novel SNP, C2136A, was detected with the HRM in the domain III (Fig. 4C and D) resulting in a Phe to Leu change in the position 712. When this mutation simultaneously occurs with the T2134A mutation (Fig. 5D), the amino acid change was Phe to Ile (Fig. 2B). Seven ticks carry the C2136A mutation, two in homozygosis, and always associated with mutations in the domain II (T170C and C190A). One tick was heterozygous for a silent mutation, C2130T, that was detected in association with T170C, G215T and C2136A (Fig. 2). Ten ticks presented both G184C and T2134A SNPs simultaneously. Nine ticks carry at least one C190A allele, six in homozygosis. The HRM allowed the detection of the T170C (Fig. 3G and H), that results in a Met to a Thr change at the amino acid residue 57 in the domain II of the para-sodium channel gene of *R. microplus*. Three ticks homozygous for this mutation and one was heterozygous. Two
ticks carry this mutation associated with the synonymous SNP C189A and the mutation C190A simultaneously (Fig. 5C). The more part of the ticks presented one (41%) or two mutant loci (31%) in the para-sodium channel. Two ticks presented three SNPs and three ticks presented four SNPs at the same time. The most frequent SNP was the C190A, found in 44.8% of the ticks, followed by T2134A, found in 37.9% of the samples. All the previously known and novel mutations in the para-sodium channel were detected in single ticks with two HRM reactions, aimed at the domains II and III (Figs. 3–5).

4. Discussion

Target site insensitivity is a resistance mechanism in *R. microplus* against pyrethroid acaricides, which is conferred by one or more mutations present in the target site, the para-sodium channel (Guerrero et al., 2014). Based on the results presented here, we successfully developed two qPCR-HRM assays to genotype the G184C, C190A and T2134A mutations, associated with resistance against four compounds of the synthetic pyrethroid class of acaricides: permethrin, cypermethrin, deltamethrin and flumethrin. We were also able to detect multiple mutations and their different combinations in two side-by-side reactions (domains II and III). Genotyping of mutations directly related to acaricide resistance could provide a useful surveillance tool to monitor resistance status promptly and accurately among *R. microplus* populations. This approach could also help refine integrated tick management strategies by adapting the selection of acaricides for effective and rational *R. microplus* control based on resistance genetic data obtained in real time.

In HRM analysis, a small region of DNA (70–100 bp) spanning the SNPs of interest is amplified by PCR in the presence of a dsDNA fluorescent dye (EVA green). This dye is used at a high concentration to achieve maximum saturation of the resulting dsDNA fragment (Bass et al., 2007). A high-resolution melt step using equipment with high thermal and optical precision is then performed in order to determine the melting temperature (T_m) of the amplicon. While the dsDNA dissociates into single strands, the dye is released and the fluorescence decreases giving a melt curve profile characteristic of the amplicon sequence (Liew et al., 2004).

The C190A, T170C and C2136A mutations are predicted to cause a relatively large change in the T_m of the amplicon (Fig. 3A, G and 4C). In contrast, the G184C and the T2134A are predicted to cause a very small change in T_m in the melt curve making it more difficult to detect in the normalized curves (Figs. 3C and 4A). However, the observation of the differences in melt plots helps in distinguishing the genotypes (3D and 4B). The combinations of different mutations (C189A/C190A; C190A/G215T; T170C/G215T; T170C/C189A/C190A; T2134A/C2136C; C2130T/C2136T) are predicted to cause even larger variations in the T_m of the melt curves (Figs. 3A–5D). Using samples with known genotypes (as determined by sequencing) the developed assays were able to efficiently distinguish the different genotypes, in homozygosity and heterozygosity.

The Santa Luiza strain has been studied by synergistic bioassays and genetic studies to determine its mechanisms of resistance to permethrin (Li et al. 2007, 2008). The present study clarifies the major mechanism...
involved, which is the presence of one, two or three mutations in different amino acid residues of the para-sodium channel gene. Stone et al. (2014) made the first description of the presence of simultaneous mutations in the para-sodium channel gene in R. microplus populations from Mexico, where a combination of the M57T amino acid change with L64I or F712I was linked to high levels of pyrethroid resistance. Nevertheless, the mutations in the Santa Luiza ticks were mostly heterozygous (82.6%; Table 3). The cause for the lower resistance level in Santa Luiza is unknown when compared to the El Zamora and Yucatan strains.

El Zamora ticks presented mutations at the nucleotide positions 189 and 190 (kdr). Twenty-five percent of the ticks were heterozygous for the C190A mutation, and the remaining were homozygous. The C189A substitution is synonymous and was never found by itself and always in heterozygosity. The same observation was made by Stone et al. (2014) in screening populations from Mexico and the USA. These authors pointed out that the presence of this SNP could prevent PCR amplification of fragments using specific primers designed to detect the C190A mutation. The HRM domain II assay we developed was successful in detecting both polymorphisms and no amplification failures were detected. The Yucatan strain was the most resistant strain evaluated in this study (mean mortality between 0 and 2.85% depending on the acaricide tested). All the ticks genotyped were homozygous for the G62R and F712I mutations, which can explain the low mortality levels found and the knockdown resistance phenotype observed (Table 2, Fig. 1).

![Fig. 5. Fluorescence difference curves obtained with HRM assays for pyrethroid resistance with multiple mutations detected in the domains II and III of para-sodium channel gene of R. microplus. 5A: G184C, C190A, and C189A, C190A; 5B: C190A, G215T, and T170C, G215T; 5C: T170C, C189A, C190A; 5D: T2134A, C2136A, and C2130T, C2136A.](image)

Table 3

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Amino acid sequence</th>
<th>SNP</th>
<th>Genotype frequency (%)</th>
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<tr>
<td>Wild type</td>
<td>GG/LL/FF</td>
<td>G184C</td>
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<td></td>
<td></td>
<td>C190A</td>
<td>0</td>
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<tr>
<td></td>
<td></td>
<td>T2134A</td>
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<td></td>
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<td></td>
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<td>St. Luiza</td>
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<td></td>
<td></td>
<td>El Zamora</td>
<td>0</td>
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<td></td>
<td>Yucatan</td>
<td>0</td>
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<tr>
<td>Heterozygous</td>
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<td>T2134A</td>
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<td></td>
<td>Yucatan</td>
<td>100</td>
</tr>
</tbody>
</table>

G: Glycine; L: Leucine; F: Phenylalanine; R: Arginine; I: Isoleucine; SNP: single-nucleotide polymorphism.
This assay could also be used to mitigate the risk associated with acaricide-resistant *R. microplus* that may infest cattle presented at ports of entry by Mexico intended for export to the U.S. (González and Hernández, 2012). Among the ticks sampled at the USA Ports-of-Entry, we found a high frequency of individuals carrying mutations in para-sodium channel gene. Most of them, with C190A or T2134A and combinations of the different mutations. The presence of multiple mutations in the para-sodium channel gene is correlated with high levels of pyrethroid resistance, and here we found ticks carrying three or four SNPs in both domains II and III (Supplementary Table). More information is required about acaricide usage or resistance status of the field samples to determine if that correlation causes high levels of pyrethroid resistance as suggested by the results of the HRM analysis.

Comparing different techniques for detection of knockdown resistance mutations in mosquitoes, Bass et al. (2007) observed that HRM analysis presented a higher failure rate than other assays, such as AS-PCR and ‘Taqman’ probes. The authors suggest that the quality and quantity of DNA could be affecting the amplification, preventing the attainment of a high signal plateau in the PCR phase, which could result in inconclusive or low resolution HRM data. In our experiments we were able to obtain satisfactory amplification and discrimination among genotypes from the tick colonies as well as from the field samples. The quality and quantity of DNA obtained with the phenol-chloroform purification was high (260/280 ratio ~1.8, over 200 ng/µL of dsDNA per sample), which likely had a positive impact on the quality of our amplification.

The main advantage of the HRM, comparing to AS-PCR and Taqman is its capacity of screening different variants in a given gene that would be undetectable using allele specific probes and primers, as virtually any different nucleotide can be detected in a sequence using the high-resolution analysis of the melting temperatures of those fragments. This feature is particularly interesting to define samples to be analyzed by sequencing in SNP discovery studies. When investigating the field samples from Mexico, in most of the cases, we detected the previously known SNPs (C190A, 44.83%, and T2134A, 37.93%). However, we were able to detect novel melt curves (Fig. 3G, H, 4C, 4D, 5B, 5C, and 5D), indicating the existence of different genotypes that were later confirmed by cloning and sequencing. Using the screening approach, we were able to detect for the first time in the Neotropical region, the mutation G215T, associated with flumethrin resistance in Australia (Jonsson et al., 2010) and a novel SNP, C2136A, located in the same codon for the Phe at the position 712. However, when this mutation is present in the absence of the T2134A mutation, the amino acid is substituted for an Ile. We hypothesize that the physiological effect of this substitution is the same as the Phe to Leu, which results in resistance to pyrethroids (He et al., 1999). While there is no information about the phenotype of the different patterns of mutations regarding the susceptibility to pyrethroids among the field samples, further research is needed to test our hypothesis.

This study provides a technique that can be used for the surveillance of pyrethroid resistance in *R. microplus* populations. Our findings can be used to develop a high-throughput method to genotype and detect allele-specific mutations in *R. microplus* populations that cause outbreaks in the U.S. The rapid turnaround of results based on a high-throughput HRM acaricide resistance assay could help Cattle Fever Tick Eradication Program personnel manage the response to *R. microplus* outbreaks, and also inform decisions regarding the concern with cattle presented at ports of entry by Mexico with the intention to be exported to the U.S. that may be infested with *R. microplus* resistant to acaricides (Pérez de León et al., 2013). This approach could also be adapted to other acaricide targets and species of ticks of medical and veterinary importance. The HRM for pyrethroid resistance SNPs could also be used in integrated control programs in other parts of the world where ticks and tick-borne diseases burden the health of humans, domestic animals, and
wildlife.

5. Conclusion
A quantitative PCR-based HRM assay method was developed that detects T710C, G184C, C190A, G215T, T2134A and C2136A SNPs and their combinations in the para-sodium ion channel gene of *R. microplus*. Bioassays confirmed the existence of broad spectrum pyrethroid resistance correlated with the frequency of the mutations in the tick strains evaluated. This assay provides a useful methodology to screen mutations associated with susceptibility to pyrethroids in tick populations and thus facilitates surveillance for acaricide resistance.

Author contribution statement
G.M.K., R.J.M., J.P.T. and A.A.P.L., oversaw the research and planned the experiments. G.M.K. and D.S. executed the bioassays, PCR, cloning and sequencing; G.M.K. and J.P.T. analyzed the data and prepared figures. G.M.K., J.P.T., D.S., R.J.M., T.P.F., D.B.T. and A.A.P.L prepared and reviewed the manuscript.

Declarations of interest
None.

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Appendix A. Supplementary data
Supplementary data to this article can be found online at https://doi.org/10.1016/j.ijpddr.2019.03.001.

References


