Differentially expressed genes in response to amitraz treatment suggests a proposed model of resistance to amitraz in R. decoloratus ticks

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Keywords: Rhipicephalus decoloratus, Amitraz, Resistance, RNA-sequencing, α2-adrenoceptor, NMDA receptor, Calcium signalling

Abstract

The widespread geographical distribution of Rhipicephalus decoloratus in southern Africa and its ability to transmit the pathogens causing redwater, gallsickness and spirochaetosis in cattle makes this hematophagous ectoparasite of economic importance. In South Africa, the most commonly used chemical acaricides to control tick populations are pyrethroids and amitraz. The current amitraz resistance mechanism described in R. microplus, from South Africa and Australia, involves mutations in the octopamine receptor, but it is unlikely that this will be the only contributing factor to mediate resistance. Therefore, in this study we aimed to gain insight into the more complex mechanism(s) underlying amitraz resistance in R. decoloratus using RNA-sequencing. Differentially expressed genes (DEGs) were identified when comparing amitraz susceptible and resistant ticks in the presence of amitraz while fed on bovine hosts. The most significant DEGs were further analysed using several annotation tools. The predicted annotations from these genes, as well as KEGG pathways potentially point towards a relationship between the α-adrenergic-like octopamine receptor and ionotropic glutamate receptors in establishing amitraz resistance. All genes with KEGG pathway annotations were further validated using RT-qPCR across all life stages of the tick. In susceptible ticks, the proposed model is that in the presence of amitraz, there is inhibition of Ca2+ entry into cells and subsequent membrane hyperpolarization which prevents the release of neurotransmitters. In resistant ticks, we hypothesize that this is overcome by ionotropic glutamate receptors (NMDA and AMPA) to enhance synaptic transmission and plasticity in the presence of neurosteroids. Activation of NMDA receptors initiates long term potentiation (LTP) which may allow the ticks to respond more rapidly and with less stimulus when exposed to amitraz in future. Overactivation of the NMDA receptor and excitotoxicity is attenuated by the estrone, NAD+ and ATP hydrolysing enzymes. This proposed pathway paves the way for future studies on understanding amitraz resistance and should be validated using in vivo activity assays (through the use of inhibitors or antagonists) in combination with metabolome analyses.

1. Introduction

Rhipicephalus microplus and R. decoloratus are hematophagous ectoparasites of economic importance due to their ability to transmit several tick-borne diseases that are detrimental to the livestock and agricultural industry on a global scale (Jongejan and Uilenberg, 2004; Walker et al., 2003). Rhipicephalus decoloratus in particular poses a threat to southern Africa due to its widespread geographical distribution and its ability to transmit Babesia bigemina (causative agent of redwater), Anaplasma marginale (causative agent of gallsickness) and Borrelia theileri (causative agent of spirochaetosis) in cattle (Walker et al., 2003). The principal control strategy implemented to regulate tick populations is through the use of chemical acaricides. In South Africa, the most commonly used chemical acaricides are amitraz and pyrethroids (Baron et al., 2015; Robbertse et al., 2016; van Wyk et al., 2016).

Resistance has been reported globally against all major classes of acaricides and poses an emergent problem in the future control of ticks and their associated tick-borne diseases (Abbas et al., 2014). Resistance to acaricides can arise through several potential resistance mechanisms including penetration resistance, target site insensitivity and enhanced metabolic detoxification. Currently, acaricide resistance mechanisms in R. microplus are well documented in comparison to R. decoloratus ticks where data is lacking (Guerrero et al., 2012). Amitraz resistance in R.
like receptors in this study. An amitraz resistant strain of R. microplus has been detected in multiple tick populations across the world (Chevillon et al., 2007; Mendes et al., 2013; Soberanes et al., 2002) and the resistance mechanism has been proposed to involve metabolic detoxification mediated by glutathione-S-transferase (Guerrero et al., 2012) or single nucleotide polymorphisms (SNPs) (Baron et al., 2015; Corley et al., 2013). Recent reports have also suggested the involvement of ATP-binding cassette transporters in the detoxification of amitraz in R. microplus (Koh-Tan et al., 2016; Lara et al., 2015). To date, no amitraz resistance mechanism has been suggested for R. decoloratus ticks.

The target site of amitraz is proposed to be monoamine oxidase or the octopamine receptor (Guerrero et al., 2012; Jonsson and Hope, 2007), with a number of mutations reported to date in the octopamine/tyramine (OCT/Tyr) receptor (Baron et al., 2015; Chen et al., 2007) as well as the β-adrenergic-like octopamine receptor (βAO) (Corley et al., 2013) for R. microplus. Recent studies have also shown the presence of a new BAOR gene present in an acaricide resistant tick population along with increased expression levels of ATP-binding cassette transporters (Koh-Tan et al., 2016). As such, it appears as though different strains of R. microplus from varying geographical locations may display different amitraz resistance mechanisms (Baron et al., 2015; Corley et al., 2013; Koh-Tan et al., 2016). Screening of the R. decoloratus OCT/Tyr receptor gene revealed that the two SNPs associated with a resistant phenotype in R. microplus were not present in R. decoloratus (Baron, 2017). Previous studies also showed that recombination events in the OCT/Tyr receptor was a determinant in the selection of amitraz resistance associated alleles in R. microplus (Baron et al., 2015). Due to the lack of recombination events in the R. decoloratus OCT/Tyr receptor gene and the presence of various species-specific SNPs (Baron, 2017), an alternative amitraz resistance mechanism might be involved.

In vitro studies have shown that amitraz has the potential to inhibit monoamine oxidase as well as increase plasma glucose levels and suppress insulin concentrations in humans (Ellenhorn et al., 1997). A study conducted on mice also showed that subcutaneous injections of amitraz acted as agonists for the α2-adrenergic receptors (Hsu and Lu, 1984). Previous studies in both honeybees and mammals furthermore suggested the involvement of α2-adrenergic receptors in response to amitraz (M’diaie and Bounias, 1991; Shin and Hsu, 1994). Due to the uncertainty of which resistance mechanism(s) R. decoloratus employs against amitraz selection pressure, all possibilities need to be considered.

RNA-sequencing (RNA-seq) has the ability to provide both quantitative and qualitative information of transcripts in eukaryotes making it a powerful tool for downstream analyses (Wang et al., 2009). Recent advances in high through-put RNA-seq have opened up several avenues for investigating the mode of action of drugs and resistance (Wacker et al., 2012). Examples include the use of RNA-seq to uncover the complex nature of pesticide resistance in the bed bug, Cimex lectularius (Mamidala et al., 2012) and to determine the procurement of drug resistance in Candida albicans (Dhamgaye et al., 2012).

In this study, we tested whether the amitraz resistance mechanism in R. decoloratus involves alternative metabolic mechanisms compared to what is currently known for R. microplus. Based on previous findings in other organisms, we hypothesize the involvement of α-adrenergic like receptors in this study. An amitraz resistant R. decoloratus strain was reared at ClinVet International (Bloemfontein, South Africa) for three generations, with amitraz selection pressure applied at every life stage. Total RNA-sequencing was performed on amitraz resistant and susceptible R. decoloratus samples, followed by de novo transcriptome assembly and subsequent analysis and annotation of differentially expressed genes.

2. Materials and methods

2.1. Establishment of an amitraz resistant tick strain

Engorged R. decoloratus females were collected from a farm in the Coombs district near Grahamstown, Eastern Cape, South Africa.Ticks from this farm displayed various levels of amitraz resistance depending on which paddock they were collected from. Individual engorged females were placed in Petri dishes and incubated at 25–28°C with a relative humidity of 75% (Cen et al., 1998). Ovipositioning occurred 7–18 days after collection and eggs from each female were then placed into separate glass vials. Larvae hatched 30–40 days after initial collection of engorged females. After 16–21 days of the larvae hatching, they were used in the Shaw Larval Immersion Test (SLIT) (Shaw, 1966).

All SLITs were completed at the Pesticide Resistance Testing Facility (PRTF, University of Free State in Bloemfontein, South Africa) in collaboration with Ms Ellenie van Dalen. Briefly, larvae were treated with 250 ppm amitraz which represents the normal field concentration and two times the LC99 value of amitraz to susceptible tick populations. Water was used as both the diluent for preparing the amitraz concentration and as a control. Larvae were placed onto one side of a circular (120 mm diameter) Whatman no 1 filter paper, with a second filter paper placed on the larvae to form a sandwich and the amitraz poured onto the filter paper. Larvae were exposed to amitraz for 10 min, after which the filter paper was removed from the amitraz solution, opened and placed onto a larger dry piece of filter paper. Using a paintbrush, larvae were stroked into filter paper envelopes, closed and incubated in humidity containers (RH > 70%). After 72 h, the envelopes were removed from the incubators and the percentage mortality of the larvae calculated.

Larvae that survived an amitraz concentration of 250 ppm were considered resistant, and were sent to ClinVet International (Bloemfontein, South Africa) to be reared on Holstein-Friesian cattle. TICKS were cycled for three generations under amitraz selection pressure which included amitraz dip exposure (250 ppm) at every developmental stage across all three generations. Tick samples from the second generation were used for RNA-sequencing while those from the third generation were used for qPCR validation studies. Due to low tick numbers in the second generation, only amitraz resistant nymphs could be collected before amitraz exposure (t = 0) and 4 h after amitraz exposure (t = 4) for RNA-seq. Adequate tick numbers in the third generation allowed for the collection of all three life stages before amitraz exposure (t = 0) and the collection of nymphs and adults 4 h after amitraz exposure (t = 4). Due to the size of the larvae, collections 4 h after amitraz exposure could not be performed for this life stage. An amitraz susceptible strain obtained from ClinVet International was used as the control. Tick samples collected were homogenized in TRI Reagent® (Sigma-Aldrich) and frozen in liquid nitrogen and stored at −80°C until RNA isolation was performed.

2.2. RNA isolation and sequencing

Total RNA isolation was performed on all samples using phenol-chloroform extraction followed by the RNeasy Mini Kit (QIAGEN®, USA). Additionally, 30 U of DNase1 suspended in RDD buffer (QIAGEN®, USA) was added to the column membrane before the final wash steps. The purified RNA was eluted in 50 μl RNase-free water. The quality of isolated RNA was evaluated using the Nanodrop-1000 spectrophotometer (Thermo Fisher Scientific, USA) and the Agilent 2100 Bioanalyzer (Agilent Technologies, USA). RNA samples which had a RNA Quality Indicator (RQI) value of 10 were sent for RNA-sequencing. All biological replicates for one treatment condition were pooled together and sequenced at BGI, Hong Kong. Three sequencing lanes were used for the three different treatment conditions (susceptible non-treated control, resistant non-treated t = 0, and resistant treated t = 4). RNA-sequencing was performed (proprietary of BGI) as follows; the
total RNA was treated with DNase I and then enriched for mRNA using oligo (dT) magnetic beads. The mRNA was fragmented using a fragmentation buffer followed by cDNA synthesis, size selection and PCR amplification. Sequencing was conducted on the Illumina-HiSeq™ 2000 next generation sequencing platform.

### 2.3. De novo assembly of sequence data and validation

All reads were first analysed, filtered and trimmed by removing the first 12 bp from each read, eliminating sequence errors and removing low quality bases using FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and ConDeTri (Smeds and Kunstner, 2011). Velvet v 1.2.10 (Zerbino and Birney, 2008) and Oases v 0.2.08 (Schulz et al., 2012) were used to perform the de novo assembly of all reads for each individual sample representing the different treatment conditions. Different k-mer values were tested during the assembly ranging from 30 to 60 k-mer in increments of 5. The Oases merge function was used to further merge and process Velvet assemblies. All transcripts less than 200 bp were removed, and a representative transcriptome for each individual sample representing the different treatment conditions.

### 2.4. FPKM and logFC determination

Bowtie (Langmead et al., 2009) was used to perform the alignments of Illumina reads for R. decoloratus nympha libraries to the de novo assembled representative transcriptome. The eXpress package v 1.5.1 was then used to estimate fragments per kilobase of transcript per million mapped reads (FPKM) values. In addition, edgeR v 3.4.0 was subsequently used to calculate a different parameter for differential gene expression, log fold change (logFC) and its corresponding p-value (Robinson et al., 2010). In all the latter, the standard settings were used for each of the software. The log fold change (FC) was plotted against −log10 p-values to obtain a volcano plot for all expressed transcripts for all treatment condition comparisons. Not all differentially expressed transcripts were significant and were further filtered based on a false discovery rate (FDR) value of 0.01, differential expression fold change and a significant P-value < 0.01.

### 2.5. Annotation of significant differentially expressed genes

Due to the very large transcriptome assemblies, only the top significant differentially expressed genes were functionally annotated. Annotations were performed using an automatic functional annotation and classification tool (AutoFACT) v 3.4 (Koski et al., 2005) at the CCG, as well as BLAST2GO v 4.1.7 (Conesa et al., 2005) at the University of Pretoria, South Africa. Blast searches were performed against the full NCBI non-redundant databases with an e-value threshold of 1e−05 to ensure confidence in annotation. Additional functional annotation included GO terms and KEGG pathways while protein function classification was performed using InterProScan (Jones et al., 2014).

### 2.6. Validation using RT-qPCR

Primers were designed using Oligo™ 7 Primer Analysis Software and obtained from Whitehead Scientific (South Africa). Primers were 21 bp in length, had a GC content between 50 and 60%, a melting temperature of 56–58 °C with no hairpin loops or primer-dimer formations predicted. Primer information and amplicon lengths can be found in Table 1 along with reference genes published by Nijhof et al. (2009) used in the study. The reference genes used were defined by Nijhof et al. (2009) and were therefore used for normalization of all expression data.

As stipulated previously, the quality of the RNA samples was analysed using the Agilent 2100 Bioanalyzer (Agilent Technologies, USA). Samples had to have an RQI value of ≥ 9 to be used for RT-qPCR. For the amitraz susceptible control, there were four biological controls for each life stage (12 samples in total). These controls represented the baseline for expression of genes predicted to be associated with amitraz resistance. Four biological controls for each life stage (12 samples in total) was available for amitraz resistant samples prior to treatment (t = 0). Four biological controls were available for only nymph and adult stages (8 samples in total) for amitraz resistant samples 4 h after treatment (t = 4), as the treated unfed larvae were too small for collection from the calves. The biological replicates used for RNA-sequencing were also used for RT-qPCR analysis (not the pooled samples) and formed part of the samples mentioned above.

Synthesis of cDNA was performed from 2 μg from each sample of RNA using the SuperScript® VILO™ cDNA synthesis kit (Thermo Fisher Scientific, USA). Briefly, a 20 μl reaction was prepared by adding the SX VILO reaction mix, 10X Superscript enzyme, RNA (2 μg) and RNase free water. The reaction is then incubated at 25 °C for 10 min, 42 °C for 60 min with a final termination step of 85 °C for 5 min.

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### Table 1

| Reference Gene | Forward Primer 5′-3′ | % GC content | Reverse primer 5′-3′ | % GC content |
|----------------|----------------------|--------------|----------------------|--------------|
| ELF1α          | CGTCTCAAGAATGGGTGCGT 45.5 | CTCAGTGGTCAGGTTGGCAG 60.0 |
| PPIA           | CGGAGGCGATTAGTAAATGAGC 50.0 | ATGAAAGGTGGGAGATGAGC 52.6 |
| ACTB           | CCCATGCAGGAGCTTCAGTAG 57.1 | CGGATCATACGAACAGGGG 57.1 |
| RLP4           | AGTTCCCTCCCGAGGGTCGAG 63.1 | GTCCTCTATACCTCTCCCG 52.4 |

* ELF1α is elongation factor 1-alpha which is a component of the eukaryotic translational apparatus. PPIA is cyclophilin which facilitates protein folding. ACTB represents beta actin which is a cytoskeletal structural protein. RLP4 is ribosomal protein L4 which is a structural component of the 60S ribosomal subunit.
The efficiency of all PCR reactions was tested prior to RT-qPCR analysis. As a general test, template amplifications were performed with 5 μM primer and 20 ng/μl cDNA samples in 25 μl reactions using EconoTaq PLUS GREEN 2X Master Mix (Lucigen, USA). Amplification reactions were run on a 2% (w/v) agarose gel to ensure efficient amplification. All primer amplification temperatures were optimized ranging between 53 and 55 °C.

The QuantStudio 12K-flex system was used for all reactions in a 384-well plate with its corresponding software (Thermo Fisher Scientific, USA). All RT-qPCR reactions were performed using the KAPA SYBR® FAST qPCR Kit Master Mix (2X) Universal (KapaBiosystems, USA). Briefly, 10 μl reactions were set up by adding 5 μl KAPA SYBR FAST qPCR Master Mix (2X), 200 nM final concentration of the forward primer and reverse primer and PCR-grade water. All reactions were done in triplicate to improve statistical analysis.

A standard curve was set up to determine the efficiency of all primers (gene targets and reference genes). Reactions for the standard curve plate were in triplicate for five dilution factors of cDNA samples (1/5, 1/10, 1/20, 1/50 and 1/100). A no template control and no transcriptase control were included for each primer set as well. Analysis of the standard curve was performed using qbase + software. The R² of the curve had to be more than 0.99 to provide good confidence within the correlation. The remaining reactions were run on the QuantStudio 12k-flex system and analysed using the qbase + software. These reactions were set up in triplicate along with no template controls and no transcriptase controls for each gene. The sample maximization method was used when setting up the plates, this reduced technical variation between samples and did not require inter-run calibration. Four reference genes (Table 1) were used for normalization of expression data.

3. Results

3.1. RNA-sequencing results

The quality of the sequencing data is shown in Table 2. The results indicate that after the filtering and trimming of raw data, the remaining reads are of very good quality (> 99%). Raw reads were submitted to NCBI (SRA accession no: SRP137618).

3.2. De novo assembly validation

The amitraz susceptible and amitraz resistant (t = 0 and t = 4) nymph samples were de novo assembled individually and then combined into one representative transcriptome using cd-hit (Li and Godzik, 2006) for R. decoloratus nymphs. The summary statistics for the individually assembled transcriptomes are shown in Table 3, and that of the representative transcriptome in Fig. 1.

After combining these transcriptomes into a representative transcriptome (Fig. 1), the final contig number was 84,871 with a mean contig length of 1235 bp and a maximum length of 18,493 bp. The de novo assembled representative transcriptome was analysed using the arthropod gene sets in the BUSCO database (Simão et al., 2015). Results

### Table 2

| Sample | Sequencing strategy | Raw data size (bp) | Raw reads number | Clean data size (bp) | Clean reads number | Clean data rate (%) |
|--------|---------------------|--------------------|-----------------|---------------------|------------------|-------------------|
| SS     | PE100               | 6 581 668 116      | 65 818 315      | 6 526 609 041      | 65 388 748       | 99.34             |
| RR_T0  | PE100               | 6 806 329 958      | 68 064 913      | 6 779 135 611      | 67 859 478       | 99.69             |
| RR_T4  | PE100               | 6 740 178 084      | 67 402 539      | 6 710 410 459      | 67 176 992       | 99.66             |

SS represents amitraz susceptible nymphs (control), RR_T0 represents amitraz resistant nymphs prior to amitraz exposure and RR_T4 represents amitraz resistant nymphs 4 h after amitraz exposure. PE indicates paired-end sequencing.

3.3. Diﬀerentially expressed genes and annotation

Differentially expressed genes (DEGs) were identified using edgeR (Robinson et al., 2010) by comparing the nymph transcriptomes from different treatment conditions to one another. The log2FC was plotted against the −log10 p-value for all DEGs (Fig. 2). The volcano plot that was generated illustrates that the most significant DEGs occur at the top of the plot while the least significant occur at the bottom. Transcripts that were upregulated are plotted to the right (> 0) and those that were downregulated occur on the left (< 0).

DEGs were further filtered based on FDR (0.01), significant P-value < 0.01, and differential expression fold change (logFC). This resulted in 1 079, 1000 and 628 significant DEGs for comparisons between SS and RR_T4, SS and RR_T0 and RR_T0 with RR_T4 respectively. All of these transcripts (2707 in total) were analysed against the NCBI non-redundant database using AutoFACT (Koski et al., 2005) and BLAST2GO (Conesa et al., 2005). Approximately 80% of the differentially expressed transcripts were without significant BLAST hits at the pre-set minimum threshold of 1e−05. This threshold was implemented using guidelines from previous studies where tick transcriptomes were annotated (De Marco et al., 2017; Guerero et al., 2016). The most significant DEGs had no BLAST hits or functional annotations and as such remain hypothetical protein coding sequences. Assigning gene ontologies (GO)’s was problematic, where ~90% of the sequences could not be assigned GO terms. From those that could be annotated, transferases appeared to be upregulated in resistant samples which may affect certain biological processes (Fig. 3). Additionally, response to stimulus was also upregulated in resistant samples.

In addition to this, InterProScan annotations were also performed using BLAST2GO (Conesa et al., 2005). Due to the lack of information provided by BLAST, GO and InterProScan annotations, KEGG pathways were subsequently considered. Only nine transcripts could be assigned KEGG ontologies (Table 4). Seven contigs with annotations matching known enzymes were identified as upregulated in amitraz resistant samples, and two were down-regulated (Table 4). Supplementary File 1 contains the significant DEGs, logFC, p-value and FDR values as well as annotation results.

### Table 3

Summary statistics of individual de novo assembled transcriptomes for R. decoloratus nymphs under different treatment conditions.

| Representative assemblies | Number of contigs | Longest contig size | Shortest contig size | Mean contig size |
|---------------------------|-------------------|---------------------|---------------------|-----------------|
| SS                        | 68,170            | 13,860              | 201                 | 634             |
| RR_T0                     | 88,086            | 16,359              | 201                 | 1188            |
| RR_T4                     | 104,985           | 18,493              | 201                 | 1193            |

SS represents amitraz susceptible nymphs (control), RR_T0 represents amitraz resistant nymphs prior to amitraz exposure (t = 0) and RR_T4 represents amitraz resistant nymphs 4 h after amitraz exposure (t = 4).

show complete BUSCOs of 96.6% with single-copy genes (57%), duplicated genes (39.6%), fragmented genes (2.3%) and missing genes (1.1%).
3.4. Validation of expression profiles

RT-qPCR was performed on individual biological replicates in triplicate using the QuantStudio 12K-flex system (Thermo Fisher Scientific, USA). Expression profiles of all transcripts in Table 4 were assessed for all biological controls across all life stages and treatment conditions. Fig. 4 shows the expression profile of all the genes relative to the life stage and treatment condition. The calibrated normalized relative quantities (CNRQ) were calculated from all biological controls (in triplicate) to generate the average CNRQ value per life stage per transcript. Analysis showed that the comparison of expression profiles between resistant and susceptible samples was significant (P-value < 0.05) while comparisons between the two resistant conditions were not (P-value > 0.05). For this reason, although differences in expression levels between the two resistant conditions were noticed, they were considered non-significant. The most significantly expressed transcript across all the life stages when comparing susceptible versus resistant was Rde_RR_057328 (P-value 0.0007887) which resembles the glycine N-methyltransferase and Rde_RR_062143 (P-value 0.00332) which resembles a pregnenolone sulfotransferase.

The general trend observed correlates with the RNA-seq findings where genes that were downregulated in amitraz susceptible samples and upregulated in resistant samples are validated by qPCR data. Two transcripts (Rde_RR_070409 and Rde_RR_081765) that were found to be downregulated in resistant samples (Table 4) were upregulated in the susceptible samples as expected. Fig. 5 shows the overall general trend of expression for all transcripts for each treatment condition, where CNRQ averages for each life stage was determined. RNA-seq was performed on pooled samples and so logFC and FPKM values cannot be directly compared to CNRQ values for this study.

4. Discussion

Rhipicephalus decoloratus ticks occupy large geographic areas of southern Africa, and are adept in transmitting several tick-borne diseases (Walker et al., 2003). By means of Shaw Larval Immersion Test (SLIT) bioassays, several R. decoloratus populations in South Africa have been shown to be resistant to amitraz (Pesticide Resistance Testing...
detected in this study, as the most significant transcripts that could be annotated with GO terms and KEGG pathways (De Marco et al., 2017). The lack of available annotations could be attributed to the prevalence of unique genes within ticks, as comparative analysis of other tick species and arthropods revealed 10,835 unique protein coding genes (Barrero et al., 2017). Additionally, it was also shown that from 2034 ESTs from R. sanguineus, 1024 were unique (Anatriello et al., 2010). This could potentially explain the lack of annotatable genes detected in this study, as the most significant DEGs in response to amitraz may be unique to ticks or specifically to R. decoloratus.

The quality of all isolated RNA was assessed at the University of Pretoria using a spectrophotometer and Agilent Bioanalyzer. It was further assessed upon arrival at BGI prior to sequencing. RNA-sequencing results were validated using RT-qPCR techniques using the appropriate controls and reference genes. Mentionable limitations to this study include; pooling the RNA samples for sequencing, not performing test PCR amplifications on the RNA, and obtaining the susceptible control from a different genetic background as compared to the resistant samples.

It is suggested that amitraz acts as an agonist on the adrenergic signalling pathway (Hsu and Lu, 1984; Young et al., 2005). The few transcripts that could be annotated with GO terms and KEGG pathways in this study identified nine enzymes that could be successfully linked to synaptic signalling. To date, adrenergic signalling in invertebrates is better characterized than in invertebrates. In vertebrates, it has been shown that amitraz acts as a α2-adrenergic receptor agonist preventing norepinephrin (NE) release from synapse membranes in mammalians (Young et al., 2005), including mice (Hsu and Lu, 1984). An agonist, such as amitraz, will bind to the α2-adrenoceptor, which is a G protein coupled receptor found in both pre- and post-synaptic adrenergic neurons (Ma et al., 2005). When an agonist binds to the receptor it results in G-protein activation with subsequent inhibition of adenyl cyclase, phospholipase C (PLC), influx of intracellular calcium (Ca++) and increased efflux of potassium (K+) ions. This combination of events results in presynaptic membrane hyperpolarization (making the cell membrane potential more negative) which inhibits action potentials between neurons resulting in sedation, paralysis and probable death (Ma et al., 2005).

In contrast, adrenergic signalling in invertebrates is most likely accomplished through the octopaminergic system where octopamine (a NE analogue) is an important neuromodulator (Blenau and Baumann, 2001) and known to play a vital role in synaptic plasticity (Koon et al., 2011). Octopamine receptors have been classified as α- or β-adrenergic-like due to their resemblance with vertebrate receptors affecting either Ca2⁺ or cAMP levels, respectively (Balfanz et al., 2005; Pfleger and Stevenson, 2005). Agonists acting on the α1-adrenergic-like receptor tend to activate phospholipase C (PLC) resulting in an increase in intracellular Ca2⁺ levels (Evens and Maqueira, 2005). In contrast, agonists of α2-adrenergic-like receptors decrease adenyl cyclase activity as well as intracellular Ca2⁺, which in turn inhibits the release of neurotransmitters such as glutamate (Dong et al., 2008; Giovannitti et al., 2015; Pan et al., 2002). However, when an agonist binds to the β-adrenergic-like receptor, adenylate cyclase is activated and cAMP levels rise (Blenau and Baumann, 2001). If we consider that amitraz acts as an agonist of the α2-adrenoceptor in vertebrates, it could potentially invoke a similar response on α2-adrenergic-like octopamine receptors in invertebrates. Based on the results obtained from this study, there could potentially be a rescue mechanism that is present in amitraz resistant R. decoloratus ticks which involves ionotropic glutamate receptors to enhance synaptic transmission and plasticity in the presence of neurosteroids.
Table 4

| Transcript | Enzyme | E-value | Pathway | Function | logFC | FPKM | Count | LogFC | FPKM | Count |
|------------|--------|---------|---------|----------|--------|------|-------|--------|------|-------|
| Rde_RR_062143 | steroid sulfotransferase (EC:2.8.2.2) | 2.8.2.2 | steroid hormone biosynthesis | Sulfation of Preg and DHEA | 1.54E-145 | 7545 | 1.82E+00 | 26 | 7415 | 6.32E+00 | 88 |
| Rde_RR_022093 | - | 3.1.1.94 | catechol O-methyltransferase | Degrades catecholamines | 4.34E-46 | 6626 | 2.47E+00 | 13 | 6069 | 0.00E+00 | 0 |
| Rde_RR_049387 | - | 3.1.3.41 | - nitrophenyl phosphatase | Aminobenzoate degradation | 3.72E-39 | 6757 | 3.83E+00 | 15 | NA | 0.00E+00 | 0 |

# 4.1 Synaptic transmission under normal conditions (Fig. 6A)

Synaptic transmission under normal conditions involves the synthesis and uptake of neurotransmitters (such as glutamate) into the presynaptic cell through neurotransmitter transporters (Destexhe and Mainen, 1994; Levitan and Kaczmarek, 1991). Glutamate is a well-known excitatory neurotransmitter in glutamatergic synapses where it is released from the presynaptic terminal and plays a vital role in transmitting signals between nerve cells (Curtis and Watkins, 1960). The glutamate neurotransmitter is stored in synaptic vesicles, and released into the synaptic cleft through exocytosis in response to an action potential. Action potentials are generated when presynaptic voltage gated ion channels allow the influx of Ca^{2+} and Na^+ ions into the cell, as well as the efflux of K^+ ions. The influx of calcium into the cell is what allows for Ca^{2+}-mediated neurotransmitter release from the presynaptic terminal (Meldrum, 2000). Glutamate can then bind to and activate postsynaptic transmitter receptors such as postsynaptic ionotropic and metabotropic glutamate receptors (Blanke and Van Dongen, 2009; Bortolotto et al., 1999) or second messenger gated channels (Destexhe and Mainen, 1994). These receptors can then exert their effects through complex second messenger systems.

# 4.2 Synaptic transmission for amitraz resistant ticks upon exposure to amitraz (Fig. 6B)

The following mechanism is proposed for synaptic transmission in amitraz susceptible ticks. When amitraz interacts with the α2-adrenergic-like octopamine receptor (OCTR) in susceptible ticks, the influx of K^+ ions is activated leading to membrane hyperpolarization. This altered state in membrane polarity subsequently prevents the entry of Ca^{2+} ions into the presynaptic cell (Giovannitti et al., 2015; Ma et al., 2005). This diminished action potential inhibits the release of glutamate neurotransmitters and prevents synaptic transmission at the synaptic cleft (Pan et al., 2002). Hyperpolarization of the post-synaptic neuron occurs due to the influx of K^+ ions. This negative membrane potential results in a net inward force of Mg^{2+} ions that enter into the cell where the effects may result in sedation, paralysis or death (Ma et al., 2005). A strong depolarization is required to dislodge the Mg^{2+} ion in the pore, allowing the permeation of ions across the membrane (Blanke and Van Dongen, 2009).

# 4.3 Synaptic transmission for amitraz resistant ticks upon exposure to amitraz: a possible rescue mechanism (Fig. 6C)

4.3.1. Pregnenolone sulfate (PregS)

A steroid sulfotransferase (EC:2.8.2.2) coding transcript (Rde_RR_062143) was found to be upregulated in amitraz resistant ticks. This sulfotransferase mediates the conversion of pregnenolone to pregnenolone sulfate (PregS) or dehydroepiandrosterone sulfate (DHEAS). PregS and DHEA-S have been shown to function as endogenous neurotransmitters or neuromodulators (Kostakis et al., 2013) and are synthesized from cholesterol in the central nervous system (Baulieu et al., 2001). PregS and DHEA-S are thought to act through G-protein dependent pathways (ffrench-Mullen et al., 1994), that result in Ca^{2+} influx into cells and allow the release of neurotransmitters from the presynaptic terminal (Valenzuela et al., 2008). PregS can act as an agonist on calcium-permeable transient receptor potential (TRP) channels (Wagner et al., 2008) or through the direct increase in Ca^{2+} levels via a NMDA-dependant pathway (Kostakis et al., 2013). Recent studies have suggested that PregS acts presynaptically through modulation of TRP channels and promoting the insertion of ionotropic receptors to the postsynaptic neuron (Kostakis et al., 2013; Smith, 2014; Valenzuela et al., 2008).

The synthesis and secretion of PregS (or a PregS analogue) in
Amitraz resistant *R. decoloratus* ticks is therefore hypothesized to enhance Ca\(^{2+}\) influx into cells, either through modulation of TRP channels or direct interaction with NMDA receptors. This may allow Ca\(^{2+}\) mediated glutamate release which is prevented in amitraz susceptible ticks upon exposure to amitraz. Due to the importance of PregS in the re-instatement of glutamate release from the presynaptic terminal, applying an inhibitor of PregS may circumvent amitraz resistant tick populations. It has been shown that endoxifen is a potent inhibitor in the sulfation of pregnenolone and DHEA (Squirewell et al., 2014) and could be included in future bioassays to test this hypothesis.

**4.3.2. Activation of ionotropic receptors**

The PregS induced influx of Ca\(^{2+}\) ions into the presynaptic cell induces a positive action potential resulting in membrane depolarization. This will allow for the release of glutamate for the initiation of synaptic transmission. Glutamate will initially bind to the ionotropic α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptor allowing the influx of Na\(^{+}\) ions and brief membrane depolarization (Blanke and Van Dongen, 2009; Bortolotto et al., 1999; Meldrum, 2000). This transitory depolarization of the membrane is sufficient to dislodge the Mg\(^{2+}\) ion from the NMDA channel pore (Blanke and Van Dongen, 2009).

Activation of the NMDA receptor is then achieved through the dual co-agonistic binding of glutamate and glycine allowing for the influx of Ca\(^{2+}\) ions into the postsynaptic cell (Blanke and Van Dongen, 2009; Dingledine et al., 1990). As previously mentioned, glutamate is a well-known excitatory neurotransmitter while glycine can serve both inhibitory and excitatory functions in the nervous system. In a study by...
Dingledine et al. (1990), it was shown that glycine promotes the actions of glutamate on the NMDA receptor. Glycine released from the glycnergic presynaptic cell contributes a small amount of glycine at the glutamate synapse, with reuptake of the majority of glycine by glycine transporters (GlyT2) (Betz et al., 2006). It has been shown that the majority of the glycine used for the activation of NMDA receptors is generated by neighbouring astrocytes where the GlyT1 transporters regulate glycine (Betz et al., 2006). The transcript Rde_RR_057328, a putative glycine N-methyltransferase (EC:2.1.1.20) was upregulated in resistant samples. Glycine N-methyltransferase converts glycine to sarcosine which is a natural inhibitor of the GlyT1 transporter. This inhibition prevents the reuptake of glycine into the astrocyte thereby contributing to the modulatory effects on NMDA receptor activation (Long et al., 2006). Even low concentrations of ambient glycine have been shown to be able to activate the NMDA receptor in conjunction with glutamate (Blanke and Van Dongen, 2009).

Activation of postsynaptic NMDA receptors will allow the influx of both Ca\(^{2+}\) and Na\(^{+}\) ions into the cell resulting in membrane depolarization. The large influx of Ca\(^{2+}\) ions into cells triggers a cascade of intracellular processes and can result in long term potentiation (LTP). LTP allows the excitatory synapse to strengthen based on continued patterns which then produces increased signal transduction between two neurons (Malenka and Nicoll, 1999; Sweatt, 1999).

It is hypothesized that in resistant R. decoloratus ticks, glycine is released from the astrocyte with reuptake transporters inhibited by sarcosine. This action compliments the presynaptic release of glutamate as both molecules are required to activate postsynaptic NMDA receptors. This mechanism assists with activation of ionotropic receptors possibly leading to LTP. This LTP mechanism could be largely beneficial for resistant ticks as their synaptic strength increases in response to amitraz. This could potentially lead to less energy required to stimulate action potentials between neurons in the event of amitraz exposure in...
future.

4.3.3. Energy regulation

Cytochrome c oxidase (EC:1.9.3.1) was downregulated in amitraz resistant samples. Previous studies have shown that high intracellular levels of Ca²⁺ can inhibit cytochrome c oxidase activity. By down-regulating cytochrome c oxidase, the entire respiratory chain is slowed down to prevent further uptake of Ca²⁺ into the mitochondria which may be detrimental to the mitochondrial function (Vygodina et al., 2013). When the mitochondrial function becomes compromised by increased Ca²⁺ levels, ATP production through oxidative phosphorylation is hindered resulting in the need for an alternative energy source. Astrocytes and neurons have the ability to generate ATP through glycolysis as a subsidiary energy source (Liu et al., 2006). An essential cofactor, NAD⁺, is required for this reaction to take place. NADH reductase (EC:1.6.5.3) which forms NAD⁺ in the mitochondria was upregulated in amitraz resistant ticks. In response to stress, the level of NAD⁺ rises which is critical to maintain neuronal survival and protection against excitotoxicity (Liu et al., 2008). During stressful conditions, the level of ATP produced can be detrimental to neuronal survival by promoting neurodegeneration (Sperlagh et al., 2006). Nucleoside triphosphate diphosphohydrolases (NTPDases) are well known enzymes of which all represent possible new targets for improved drug resistance. In conclusion, the results from this study identify NTPDase-like enzyme was upregulated in amitraz resistant ticks and could potentially protect synapses from increased ATP levels at synaptic junctions by converting excess ATP to adenosine under stressful conditions.

4.3.4. Modulation of activation processes

Overactivation of the NMDA receptor can result in excitotoxicity and therefore requires careful modulation of its activity (Blanke and Van Dongen, 2009). In this regard, there was down-regulation of a transcript (Rde_RR_070409) resembling estrone sulfotransferase which converts estrone to estrone-3-sulfate. Estrone is known to attenuate NMDA excitotoxicity in an estrogen receptor-independent manner (Kajta et al., 2002) and potentially through antagonizing caspase-3-like mechanisms (Kajta et al., 2004). This action of estrone on the NMDA receptor possibly prevents over activation and subsequent excitotoxicity in amitraz resistant ticks. Additionally, there was upregulation of catechol O-methyltransferase (COMT), transcript Rde_RR_022093, which degrades catecholamines such as nor-epinephrine (NE) (or in this case potentially octopamine which is an analogue of NE in invertebrates). Degradation of these catecholamines is an important modulatory effect as they would continue to further agonise α2-adrenoceptors (Axelrod, 1957). These mechanisms could potentially represent neuroprotective attributes of the amitraz resistant phenotype and prevent over activation of synaptic transmission in resistant R. decoloratus ticks.

Future studies should include validating the proposed pathway through additional in vitro functional assays as these proposed mechanisms have not been well characterized in invertebrates. These enzymes should also be evaluated in the economically important tick species, R. microplus, to determine if both tick species display similar responses to amitraz exposure. Further validations should include the use of agonists/antagonists as well as metabolomics to fully validate the proposed resistance mechanism. The proposed mechanism does not exclude the involvement of other enzymes/proteins nor does it exclude the contributions of SNPs to amitraz resistance. The differentially expressed enzymes described in this model may also contribute to various other biological pathways that may/may not also contribute to amitraz resistance. In conclusion, the results from this study identified nine enzymes of which all represent possible new targets for improved drug development. The synthesis of these enzymes, their transporters and associated targets can all be considered for improved tick control strategies in the future.

Acknowledgments

Funding was provided by: (a) Gauteng Department of Agriculture and Rural Development, C Maritz-Olivier; (b) Zoetis South Africa Pty Ltd., C Maritz-Olivier; (c) National Research Foundation, THRIP grant nr: 83890, C Maritz-Olivier. Zoetis South Africa assisted with sample collection. None of the other funders had any role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript. Dr Eshchar Mizrahi for the input into the data analysis and manuscript.

Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.ijpddr.2018.06.005.

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