Responses to *Varroa destructor* and *Nosema ceranae* by several commercial strains of Australian and North American honeybees (Hymenoptera: Apidae)

Thomas E Rinderer,1* Benjamin P Oldroyd,2 Amanda M Frake,1 Lilia I de Guzman1 and Lelania Bourgeois1

1Honey Bee Breeding, Genetics, and Physiology Laboratory, USDA-ARS, 1157 Ben Hur Road, Baton Rouge, LA 70820, USA.

2Behaviour and Genetics of Social Insects Laboratory, School of Biological Sciences A12, University of Sydney, Sydney, NSW 2006, Australia.

Abstract

The potential impact of varroa (*Varroa destructor*, Anderson & Trueman) on Australian beekeeping and agriculture depends in part on the levels of resistance to this parasite expressed by Australian commercial honeybees (*Apis mellifera*). The responses of seven lines of Australian honeybees to *V. destructor* were compared with the responses of a stock of Italian honeybees from the United States known for its susceptibility to *V. destructor*, Russian honeybees (RHB) and a stock expressing the varroa sensitive hygiene trait (VSH). The experiment began in May with uniform colonies having uniform infestation of *V. destructor*. *V. destructor* infestations measured as the percentage of adult bees infested in the Australian lines and the Italian stock rose from less than 10% in August to over 25% in October. From August to November, 44% of both the Australian and Italian colonies died while strongly exhibiting symptoms of parasitic mite syndrome. In contrast, RHB and VSH colonies displayed comparative resistance to *V. destructor*. Their infestation rates rose from about 5% in August to 10% (RHB) and 14% (VSH) in October. Likely, some of this increase resulted from invasion pressure by mites from the dying Australian and Italian colonies. During the August to November period, 4.4% of the RHB and 14.3% of the VSH colonies died. In comparisons of the seven Australian lines, only nonsignificant and trivial differences were found for infestation and mortality rates. All Australian lines were highly susceptible to *V. destructor*. Additionally, evaluations of rates of *Nosema ceranae* infections were made throughout the course of the experiment. Although high levels of infection were found across all stocks and lines, no stock or line exhibited an adverse effect from *N. ceranae* infection.

Key words


INTRODUCTION

*Varroa destructor* (Anderson & Trueman 2000) is an external parasitic mite that, until recently, infested only the Eastern hive bee, *Apis cerana* (Peng et al. 1987; Oldroyd 1999). Although *V. destructor* feeds on both adult bees and brood, it persists in colonies of *A. cerana* without causing serious harm (Koeniger et al. 1983; Peng et al. 1987; Oldroyd 1999; Rath 1999). However, *V. destructor* has successfully extended its host range to the Western honeybee, *A. mellifera*, and has spread through much of this honeybee’s Old and New World ranges (Oldroyd 1999). *V. destructor* is highly virulent on *A. mellifera* as infestations result in the death of colonies within 1–2 years of being infested without beekeeper intervention (Shimanuki et al. 1994) and from 8 months to a year in areas where brood rearing is year-round (Branco et al. 1999).

Various methods have been developed to protect colonies from *V. destructor*, including biotechnical methods to several acaricides. Biotechnical methods, principally drone brood removal, are effective but time-consuming and are only suitable for use with a small number of colonies (Boot et al. 1995; Calis et al. 1999). For the most part, large scale apiculture has relied on acaricides to control the parasite. However, several problems attend the use of acaricides: mite populations have developed resistance to acaricides, honey can be contaminated if acaricides are used during a nectar flow, and the timing of nectar flows can compete with the timing of acaricide treatments necessary to effectively control the mites. Additionally, properly timed treatments will eventually contaminate the hive to damaging levels, and acaricides decrease the reproductive potential of drones (Lodesani et al. 1995; Elzen et al. 1998; de Guzman et al. 1999).
Honeybees that resist *V. destructor* are an attractive alternative to the continued use of acaricides. ‘Russian’ honeybees (RHB), a strain developed by the USDA (Rinderer et al. 2001) have several mechanisms of resistance to varroa that act in concert to provide commercially important levels of resistance (de Guzman et al. 2007). Honeybees with the ‘varroa sensitive hygiene’ (VSH) trait detect and remove varroa-infested brood (Harris et al. 2009). Some beekeepers use RHB or VSH without reliance on acaricides, while other beekeepers use only ‘soft’ chemical treatments such as thymol.

*V. destructor* has not yet reached Australia. However, considering varroa’s successful occupation of both the Old and New World ranges of *A. mellifera*, it seems likely that varroa will soon spread to Australia. Although they may have retarded the spread of varroa, Pacific Ocean isolation and regulations against honeybee importations have not protected either New Zealand (infested with varroa since 2000) or Hawaii (infested with varroa since 2007).

Considering the likelihood that *V. destructor* will eventually reach Australia, it is prudent to know the response of commercial strains of Australian *A. mellifera* to this parasitic mite. This study compared responses of a total of 240 queens from seven strains of commercial Australian honeybees and three strains of North American honeybees. The seven Australian strains have not been bred for resistance to *V. destructor*, while two of the American strains (RHB and VSH) have documented resistance to *V. destructor* (Rinderer et al. 2001; Harris et al. 2009). The third American strain is a variety of Italian honeybees that has been bred in the presence of *V. destructor* but has not been artificially selected for resistance (Rinderer et al. 2004a; de Guzman et al. 2007).

Additionally, we evaluated the rates of infestation of the colonies in the experiment by *Nosema ceranae*. This microsporidian parasite has recently invaded the honeybee colonies of both Australia and the United States. No stock has yet been found to have strong resistance to this parasite, and finding one would be highly desirable.

**MATERIALS AND METHODS**

**Honeybee stocks**

Australian queens came from seven queen breeders that were more than 1000 km from the *A. cerana* exclusion zone. Cages were marked with letters (A through G) corresponding to queen breeders. However, line identity for queens was unknown to the persons collecting data. The queens were given water everyday during shipment from queen breeders to their introduction to colonies in apiaries in Kansas, USA, with a history of good nectar and pollen availability. The shipment of queens was approved and regulated by both the Australian Quarantine and Inspection Service (AQIS) and the USDA – Animal and Plant Health Inspection Service (APHIS). These services established various pathogen- and parasite-free certification, and transportation procedures, and facilitated and assured compliance with these regulations. Experimental apiaries were quarantined in compliance with APHIS requirements.

Resistant queens (RHB = 25 and VSH = 29) and susceptible Italian queens (25) from the United States were all obtained from commercial sources. They were shipped to the USDA, Honey Bee Breeding, Genetics and Physiology Laboratory in Baton Rouge where they were re-caged without attendants, placed in queenless colonies and transported to the experimental apiaries in Kansas.

Upon arrival in Kansas, all queens were marked with acrylic paint and wing clipped to assure later identification. Paint-marking did not identify specific stocks or lines. Small (four frames of brood with adhering bees and six frames with adhering bees) queenless colonies were prepared 1 or 2 days prior to queen introduction. For each of four apiaries, colonies were randomly assigned to queens of the four groups, such that each apiary contained a proportional representation of each group. Apiaries were separated from the next nearest apiary by at least 3.6 km, and contained 54, 57, 60 and 80 colonies. Colonies were in standard 10-frame Langstroth hives placed on four-way migratory pallets. Colonies were individually marked, and the specific queen type (Australian A through G, RHB, VSH or Italian) was recorded at introduction. On 17 May 2011, queens were introduced in cages with a candy-release system. Four days after, colonies were inspected, and successful introductions as indicated by the presence of the marked queen and eggs were recorded (Table 1).

**Varroa mite inoculations**

Colonies used to produce the queenless colonies had followed a commercial schedule for acaricide treatment for the previous 12 months. They were treated 8 months prior to the beginning of the experiment with the intention of producing test colonies with low and reasonably uniform numbers of mites. Samples of adult worker bees (300–500 bees) taken from the brood nest of each of the queenless colonies detected low infestations of varroa averaging 2.65 ± 0.22 (mean ± SE) mites. In order to assure a reasonably uniform infestation, adult worker bees from highly infested colonies were pooled in a large cage. Once inside the cage, these bees were allowed to settle for 3 h to facilitate an even distribution of mites. Samples of bees from different places within the cage were then sampled to determine their infestation levels. The bees were then divided into groups that were placed in cages formed with a standard frame and eight-mesh screen. Each experimental colony was given a cage that held 1200 worker bees, with an estimated 37.3 ± 5.0 mites. The mites were able to exit the cages through the screen and infest the experimental colonies. Cages were removed from the colonies 1 month later when the caged bees had died and the mites had gained entrance to the colony.

**Colony evaluations**

Colonies were evaluated on 14 June, 12 July, 23 August, 5 October and 3 November 2011. These evaluations consisted of
first determining queen survival. Dead colonies and colonies that had no brood suitable to use for queen rearing and with laying workers were considered dead. The presence of original queens was also determined. As the goal of the experiment was to compare the response of genetically different stocks to *V. destructor* infestation, infestation and colony size data were collected only from colonies that retained their original queen. The survival and courses of varroa and Nosema infestations were not determined for colonies with supersede queens.

For surviving colonies with original queens, samples of ca. 300–500 worker bees were collected from the surfaces of two brood frames, and estimates were made of the populations of bees according to procedures commonly used for grading colonies rented for almond pollination in California (Rinderer et al. 2011). Briefly, estimates are made from counts of the number of spaces between frames filled by bees from both the top and bottom sides of hive bodies. Bee population counts were only conducted in May, August, October and November. Bee samples were frozen on dry ice for transport to the laboratory and stored frozen (at ~20°C) until processing.

**Determinations of varroa and Nosema infestation**

Fifty bees were removed from the samples for evaluations of *N. ceranae* infestation levels. The remaining bees were washed with soapy water to remove varroa mites (Rinderer et al. 2004b), then the mites and bees were counted to determine the percentage of adult bee infestation.

*N. ceranae* infestation was determined with a real-time quantitative PCR assay (Bourgeois et al. 2012). Briefly, 30 bees from each colony were dissected, and midguts were removed. Midgut tissue was pooled per colony and homogenised. Genomic DNA was extracted, then amplified with species-specific primers and probe targeted for *N. ceranae*. Detection and quantification were performed on a StepOne™ real-time PCR System (Applied Biosystems, Carlsbad, CA, USA). Protocols for FAST PCR and reagent specifications followed those described by Bourgeois et al. (2010). Modifications to the protocol were made by elimination of primers and probe reagents for *N. apis* and addition of distilled water to adjust the reaction volume to 12.5 µL. All other reagents and specifications remained the same. All samples were run in duplicate and were directly quantified by comparison with a standard curve of known levels of *N. ceranae* DNA copy number. Standards were generated from serial dilutions of a cloned PCR product of the target sequence from *N. ceranae* (Bourgeois et al. 2010). Data were converted from copy number to Nosema spores per bee based on the formula defined in Bourgeois et al. (2010).

**Data analyses**

All analyses done for the four groups of honeybees (Australian, Italian, RHB and VSH) were also conducted for comparisons of the Australian lines. As very few Australian and Italian

---

**Table 1**

<table>
<thead>
<tr>
<th>Line</th>
<th># Queens introduced</th>
<th>% Dead or queenless</th>
<th>% Dead or queenless</th>
<th>% Supersedure</th>
<th>% Dead or queenless</th>
<th>% Supersedure</th>
<th>% Dead or queenless</th>
<th>% Supersedure</th>
<th>% Dead or queenless</th>
<th>% Supersedure</th>
<th>% Dead or queenless</th>
<th>% Supersedure</th>
<th>% Dead or queenless</th>
<th>% Supersedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>28</td>
<td>76.6</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
</tr>
<tr>
<td>B</td>
<td>22</td>
<td>76.6</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
</tr>
<tr>
<td>C</td>
<td>23</td>
<td>76.6</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
</tr>
<tr>
<td>D</td>
<td>25</td>
<td>76.6</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
</tr>
<tr>
<td>E</td>
<td>24</td>
<td>76.6</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
</tr>
<tr>
<td>F</td>
<td>25</td>
<td>76.6</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
</tr>
<tr>
<td>G</td>
<td>24</td>
<td>76.6</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
<td>19.0</td>
<td>0.4</td>
<td>10.7</td>
<td>21.4</td>
</tr>
<tr>
<td>Australian</td>
<td>171</td>
<td>64.3</td>
<td>25.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
<td>16.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
<td>16.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
</tr>
<tr>
<td>Russian</td>
<td>25</td>
<td>64.3</td>
<td>25.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
<td>16.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
<td>16.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
</tr>
<tr>
<td>VSH</td>
<td>25</td>
<td>64.3</td>
<td>25.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
<td>16.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
<td>16.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
</tr>
<tr>
<td>Italian</td>
<td>25</td>
<td>64.3</td>
<td>25.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
<td>16.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
<td>16.0</td>
<td>0.0</td>
<td>16.0</td>
<td>32.1</td>
</tr>
</tbody>
</table>

Values with the same letter are not significantly different (*P* > 0.05).

© 2012 The Authors
Australian Journal of Entomology © 2012 Australian Entomological Society
colonies were surviving in November, only analyses related to survival were conducted for November. SAS Institute (2008) was used for all analyses.

Varroa infestation

Prior to analysis, data on varroa infestation were transformed with an arcsine square-root transformation to more closely approximate normality. To control for variation among the apiaries, a randomised block design was used to determine the effect of stock and observation date on infestation levels. Mean separations were determined with post-hoc t-tests (SAS Institute 2008).

Queen survival, superscedure and colony survival

Queen longevity was calculated as the number of days a queen was known to be in the colony. Ten of the 250 queens (4%) did not survive introduction and were not included in this comparison. Dates of last observation of a queen were used as mortality dates, although all queens died sometime during the observation interval following this date. Prior to analysis, data on queen survival days were transformed with a square-root transformation to better approximate normality. Proc Lifetest was run to determine if there were differences in longevity among the stocks and lines (SAS Institute 2008). A randomised block design was then used to determine the nature of the differences. The proportions of original queen survival, supersedure rates and death/queenless rates were compared among the stocks for each observation date using the Marascuillo procedure (http://www.itl.nist.gov/div898/handbook/prc/section4/prc474.htm).

Colony sizes

Prior to analysis, data on colony size were transformed with a square-root transformation to more closely approximate normality. To control for variation among the apiaries, a randomised block design was used to determine the effect of stock and observation date on infestation levels. Mean separations were determined with post-hoc t-tests.

Varroa infestation

Numbers of *V. destructor* rose in all stocks (Australian, RHB, VSH and Italian) through time (Fig. 1). There was a significant interaction (*P < 0.0001*) between honeybee stock and date of observation. Hence, infestation, defined as adult mites per 100 adult worker bees, was evaluated for each observation. All stocks began with colonies having similarly low infestation on adult bees in May (overall average = 2.63 ± 0.23%) (*F*3,711 = 2.33, *P* = 0.073). Infestation levels rose modestly and remained similar through June (3.57 ± 0.22%) and July (5.10% ± 0.26%) (*F*3,749 = 0.59, *P* = 0.623 and *F*3,769 = 1.37, *P* = 0.250, respectively). August showed an increase in varroa mite infestations in Australian (7.80% ± 0.54%) and Italian (9.51% ± 1.36%) colonies that were significantly higher than the mite infestations in RHB (4.09% ± 0.50%) and VSH (3.79% ± 0.82%) colonies (*F*3,800 = 7.06, *P* = 0.0001). By October, overall, mite infestations had more than doubled, and the differences between stocks were consistent with August data: Australian (21.28% ± 1.93%) and Italian (17.90% ± 3.30%) colonies had significantly higher levels of infestation than the RHB (10.01% ± 1.46%) colonies and Australian colonies had higher levels than the VSH (13.57% ± 2.83%) colonies (*F*3,800 = 7.06, *P* = 0.0001).

The same general trends were found in comparisons of the seven Australian lines (Fig. 2). There was an interaction between lines and observation date (*P < 0.0001*). In May, infestations were low (2.37% ± 0.24%), with some differences between the lines: lines G and E had the greatest infes-
tations, and lines A, B, D and F had the lowest (F_{6,450} = 2.23, P = 0.039). Overall, infestations rose but were similar in the seven lines in June (3.43% ± 0.25%), July (5.34% ± 0.30%) and August (7.80% ± 0.54%). Overall, by October, varroa infestation (21.52% ± 1.96%) had increased 3.2-fold. In October, differences among lines were observed: lines C, D and F had the greatest infestations, and line G had the lowest (F_{5,520} = 5.56, P < 0.0001). No colonies of line B survived until October.

**Queen survival and supersEDURE**

Lifetest results showed that the four groups (Australian, RHB, VSH and Italian) had insignificant differences in the average number of days queens survived (χ² = 7.24, P = 0.065). Italian queens had the highest average survival of 95 ± 11 days, followed by RHB (94 ± 16 days), then VSH (83 ± 14 days) and lastly, Australian queens (80 ± 4 days). Nonetheless, these insignificant trends in survival days reflect the death of many colonies or the supersEDURE of original queens after the October inspection (Table 1). In November, survival of colonies with original queens was: RHB queens (43.5%) ≥ VSH queens (28.6%) ≥ Italian queens (12.0%) ≥ Australian queens (7.9%), with the difference in the survival of RHB and Australian queens being significant (P = 0.05). Interestingly, supersEDURE rates for the four groups were similar.

There were significant differences (χ² = 16.57, P = 0.011) in the Lifetest results for queen survival among the Australian lines (Table 1). Line E had the longest survival (105 ± 12 days), followed in descending order by D (97 ± 11 days), G (96 ± 12 days), F (75 ± 11 days), C (74 ± 11 days) and A (67 ± 9 days). Queens of the B line lived the shortest period (47 ± 8 days). By October, all colonies of line B had died or superseded. Likewise, queens of lines A, C and F had poor survival compared with lines D, E and G (Table 1). By November, line E had significantly higher original queen survival rates (29%) than lines A (0%) and B (0%). Lines G (14%), D (5%), C (4%) and F (4%) were not significantly different. The supersEDURE rate did differ significantly (P < 0.05) among the lines with line C (39%) having a higher rate than line G (5%).

**Colony survival**

Colony deaths were similar for all four groups until October when varroa infestations were at high levels (Table 1). Between August and October, colony deaths in Australian and Italian lines nearly doubled. In October, death rates were: Australian colonies (51.8%) ≥ Italian colonies (40.0%) ≥ RHB colonies (30.4%) > VSH colonies (25.0%) (P = 0.05). Colony deaths continued through October and into November, especially for Australian and Italian colonies: Australian colonies (71.3%) ≥ Italian colonies (68%) > VSH colonies (39.3%) ≥ RHB colonies (34.8%) (P = 0.05).

Mortality in the colonies of the seven Australian lines was not significantly different. Line G had the highest death rate (82%), followed in descending order by D (76%), F (76%), A (75%), B (67%), E (63%) and C (61%). The supersEDURE level differed significantly (P < 0.05) among the lines with line C (39%) being higher than line G (5%).

**Colony sizes**

In the analysis of colony size data, there was an interaction between stock and observation date (F_{7,303} = 2.11, P = 0.029) (Fig. 3). In May, at the beginning of the experiment, all colonies were of similar size (10.79 ± 0.18 frames) (F_{4,412} = 1.12, P = 0.342), although RHB colonies were numerically the smallest. By August, the Australian colonies were significantly smaller (8.14 ± 0.34 frames) than the Italian colonies (9.84 ± 0.94 frames), while the RHB (9.71 ± 0.69 frames) and VSH colonies (9.29 ± 0.93 frames) were similar (F_{4,465} = 2.67, P = 0.047). In October, the RHB (8.96 ± 0.79 frames) and VSH colonies (8.13 ± 0.72 frames) were significantly larger than Australian colonies (6.74 ± 0.42 frames), and the Italian colonies (7.63 ± 1.22 frames) were similar to the other groups (F_{5,465} = 3.34, P = 0.019).

In the comparison of the seven Australian lines, there was no significant interaction between line and observation date (F_{5,163} = 1.26, P = 0.234). There was no difference in colony sizes between the lines (F_{2,228} = 1.57, P = 0.158), but there were differences between observation dates (F_{5,163} = 44.86, P < 0.0001). Colonies were largest in May (10.91 ± 0.23 frames) and smallest in October (8.13 ± 0.72 frames).
frames), then decreased in August (8.14 ± 0.34 frames) and October (6.80 ± 0.43 frames). In November, they marginally decreased (6.18 ± 0.63 frames).

**Nosema infestation**

The analysis of *N. ceranae* infection also indicated a significant interaction between stock and observation date (F_{15,616} = 2.53, *P* = 0.001) (F_{3,163} = 1.26, *P* = 0.234). In May and June, all stocks had similar and high levels of *N. ceranae* (13.04 ± 0.17 ln spores; F_{3,606} = 2.29, *P* = 0.078 and 16.74 ± 0.14 ln spores; F_{3,606} = 1.30, *P* = 0.275, respectively), although infections in June were slightly higher than in May. In July, the VSH colonies had more *N. ceranae* (14.03 ± 0.33 ln spores/bee) than the Italian (11.91 ± 0.76 ln spores/bee), Australian (11.66 ± 0.31 ln spores/bee), and RHB colonies (11.63 ± 0.94 ln spores/bee) (F_{3,606} = 3.36, *P* = 0.018). In August, the stocks had similar rates of infection (F_{3,606} = 1.15, *P* = 0.328). In October, Italian colonies had more *N. ceranae* (9.78 ± 1.60 ln spores/bee) than Australian (7.44 ± 0.56 ln spores/bee) and RHB colonies (6.06 ± 0.47 ln spores/bee), and VSH (9.84 ± 2.45 ln spores/bee) had more than RHB colonies (F_{3,606} = 3.28, *P* = 0.021).

In the analysis comparing the levels of *N. ceranae* infection among the Australian lines, lines and observation time once again interacted (F_{26,400} = 2.14, *P* = 0.001). In May, colonies of all lines have similarly high (12.83 ± 0.20 ln spores/bee) infection (F_{5,454} = 0.72, *P* = 0.636). Overall, infection rose in June but remained similar among lines (F_{5,454} = 0.20, *P* = 0.98). In July, overall infection declined and remained equal among the lines (F_{5,454} = 1.80, *P* = 0.097). In August, infections varied among the lines (F_{5,454} = 2.94, *P* = 0.008): colonies of line E were the most infected, followed by those of lines B, F, A, C and D, with colonies of line G having the lowest numbers of spores. In October, colonies of line A had significantly higher levels than colonies of lines C, D, E, F and G (F_{5,454} = 2.37, *P* = 0.029).

**DISCUSSION**

Overall, our results strongly suggest that the Australian lines and the Italian stock were similarly susceptible or intolerant to varroa and parasitic mite syndrome (PMS). Varroa infestations in the Australian lines and the Italian stock rose from rates below economic thresholds (Delaplane & Hood 1999) in August to very high levels (above 20%) by October. After August, 43.9% of the Australian colonies and 44% of the Italian colonies died. Varroa infestation was similar for Australian and Italian colonies in October. During this period, the colonies that were most heavily infested died, and the majority of the colonies that remained were so infested that they were dying and incapable of supporting additional mite population growth.

These observations contrast with the results from RHB and VSH colonies, which displayed comparative resistance to *V. destructor*. Their infestation levels in October were 10% (RHB) and 14% (VSH) that rose to about 18% for both stocks in November. Although these levels are substantially lower than those of the Australian and Italian colonies, they are unusually high for colonies that are resistant to varroa (de Guzman et al. 2007). During the August to November period, 4.4% of the RHB and 14.3% of the VSH colonies died. Doubtless, the high infestations and colony deaths experienced by the Australian and Italian colonies from October to November were conducive to producing apiaries that had heavy mite invasion pressure (Rinderer et al. 2004a). This invasion pressure probably accounted for much of the October to November increases in varroa numbers in the resistant stocks. Typically, RHB colonies in mixed apiaries with susceptible colonies have higher levels of infestation than RHB colonies that are not in mixed apiaries (Rinderer et al. 2004a).

Generally, the varroa infestation rates suggest that there were a few differences in resistance to varroa or tolerance to the viruses they transmit (Chen et al. 2005) between the Australian lines. Colonies with queens of lines A, C, D and F had both high levels of varroa infestation and high mortality, indicating a
strong susceptibility to varroa mites. While colonies of line B had low levels of varroa in August, by October, 66.7% had died, and 33.3% had superseded. These line B colonies died with lower levels of varroa. They may have been comparatively more susceptible to varroa, less tolerant of the viruses that are transmitted by varroa (Chen et al. 2005) or their survival was less owing to an unknown cause. Colonies of line G had a similar comparatively low number of varroa but had the lowest supersede rate and the highest death rate. This line suffered from susceptibility to varroa and perhaps a heightened intolerance to transmitted viruses. Line E also supported a comparatively low population of varroa mites and had a reduced supersede rate and somewhat lower death rate compared with most of the other lines. Of all the Australian lines, line E appears to be very marginally more resistant to varroa mites. However, none of the Australian lines displayed any commercially useful levels of resistance to V. destructor.

Conclusions concerning the relative susceptibility of the stocks to V. destructor infestations based on the numbers of infesting mites and colony survival are supported by colony size data. Although all stocks began the experiment in May with similar populations, by October, the Australian and Italian colonies were respectively significantly and numerically smaller. Colonies heavily infested with varroa mites typically will dwindle as they die.

All the stocks had high levels of N. ceranae infestation throughout most of the experiment. Although a few significant differences in infestation level were observed in July, nosema infestations were not unambiguously associated with differences in mortality. For example, Italian colonies had more varroa, more nosema and higher mortality than RHB colonies in November. However, although minor differences were noted in measurements of spores per bee, no stock or line showed any apparent resistance to nosema. Likewise, nosema infestation showed no apparent association with diminished colony health during the course of this experiment. A similar pattern was observed in A. m. carnica colonies in Switzerland (Dainat et al. 2012) in contrast with the results of Higes et al. (2009) in Spain. The chief causes of colony mortality in this experiment were V. destructor and, as evidenced by the widespread PMS, the viruses that it transmits.

It is widely thought that despite regulatory controls Australia will one day be invaded by V. destructor. Clearly, the current populations of A. mellifera in Australia are highly susceptible to V. destructor. Certainly, acaricides will be the first strategy to attempt to cope with V. destructor once it reaches Australia. However, the mites may be already resistant to at least some acaricides. It is highly desirable to develop and implement strategies to obtain or develop A. mellifera that is resistant to V. destructor for Australian apiculture prior to the parasite’s arrival.

ACKNOWLEDGEMENTS

We thank T. Stelzer, J. Wagnitz, S. O’Brien, J. Wales, L. Beaman, G. Delatte, R. Watts, W. Pope, R. King and S. Tanh from Baton Rouge and Daniel Martin, Apiary Inspector, State of Victoria, Australia for their technical help, and Jerry Brown for providing the honeybee colonies in the study. We thank the staff of the USDA APHIS, and the AQIS for their work in defining stock importation procedures and facilitating the stock importation from Australia. We particularly thank the staff of Qantas Airways for facilitating stock importation. This study was funded by the Australian Government Rural Industries Research and Development Corporation.

REFERENCES


Accepted for publication 9 October 2012.