RAPD identification of *Varroa destructor* genotypes in Brazil and other regions of the Americas

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**ABSTRACT.** The mite *Varroa destructor* is the main pest causing damage to apiculture worldwide. In Brazil and other parts of the world, where bees of African origin and their hybrids predominate, the bees can survive these mites without treatment. Studies have shown a correlation between the various genotypes of the mite and its fertility in different geographical regions. Information about mite genotype could be helpful in understanding the diverse effects and relationships of the mite with bees in different regions of the world. DNA analysis by RAPD technique has permitted identification of three distinct genotypes in the mite *V. destructor*, namely Russian, Japanese and Papua New Guinea. We found predominance of the Russian genotype in Brazil, along with other parts
of South America, and in Cuba and Mexico. The Japanese genotype was exclusively found on Fernando de Noronha Island in Brazil.

Key words: RAPD; Varroa destructor; Resistance; Genotype

INTRODUCTION

Varroa destructor mite, an ectoparasite of Apis cerana and Apis mellifera honey bees, is the main pest responsible for problems in apiculture. However, the intensity of damage caused by Varroa mites has been shown to vary according to the region studied. In Brazil and other parts of the world, where bees of African origin and their hybrids predominate, a perfect relationship exists between the parasite and its host. However, it is unknown whether the severity of the effects caused by the Varroa parasite depends on the genotype of the bees and/or on the genotype of the mite. Evidence suggests that the mite V. destructor is a complex species. Studies have shown a correlation between the various genotypes of the mite and its fertility in different geographical regions (de Guzman and Rinderer, 1999). This information could be helpful in understanding the diverse effects and relationships of the mite with bees in different regions of the world.

DNA analysis by the randomly amplified polymorphic DNA (RAPD) technique has permitted a more precise study of the genetic variability among mite populations (Kraus and Hunt, 1995), and has led to the identification of three distinct genotypes of the mite V. destructor: Russian (R), Japanese (J) and Papua New Guinea (PNG) (de Guzman et al., 1997; Anderson and Fuchs, 1998). In a previous study, Kraus and Hunt (1995), using the RAPD technique, identified bands that are common to mites from Germany and the United States, but are absent in mites from Malaysia. Using the OPE-07 primer, the R genotype has been characterized by the presence of a 766-bp band, which is absent in mites with the J genotype (de Guzman and Rinderer, 1999). The use of the OPP-03 primer permitted the detection of a 442-bp band characterizing the R genotype. Genotype J does not possess this band but produces two distinct bands of 675 and 412 bp, which are absent in the R genotype.

Worldwide differences in the infestation of A. mellifera bees with the mite V. destructor suggest a correlation between some genotypes and a higher or lower virulence of the mite. DNA analyses have revealed variable V. destructor genotypes infesting A. mellifera and A. cerana in Asia (de Guzman et al., 1997; Anderson and Fuchs, 1998). A low virulence of the PNG genotype has been well documented in Papua New Guinea and Indonesia (Anderson and Fuchs, 1998). In contrast, marked losses of A. mellifera colonies are observed in countries in which the R genotype occurs, such as the Philippines and Korea (Cervancia, 1993). In Vietnam, the mite V. destructor represents the main agent causing damage to apiculture, with A. mellifera colonies harboring mites with the R genotype, while A. cerana colonies harbor mites with the J genotype (de Guzman and Rinderer, 1998). This finding may be explained by the differential reproduction of the mite in A. mellifera and A. cerana colonies, as observed in Japan, where only the J genotype (de Guzman et al., 1997) and the two bee species exist (A. mellifera and A. cerana) and where no loss of colonies has been observed since 1970 (Yoshida, cited in de Guzman and Rinderer, 1999).

Studies showed a worldwide distribution of the R genotype, including Europe, United States, Mexico, and Argentina, while the J genotype is restricted to Japan, Brazil, Puerto Rico, and some states in the United States (de Guzman et al., 1997). However, in the specific case of Brazil, analyses have been based on mites sampled only from apiaries in Rio
de Janeiro State in 1990 and 1993. Therefore, the study of mites from various other regions of Brazil, mainly the south region (Rio Grande do Sul and Santa Catarina), could reveal the presence of the R genotype also in Brazil, especially considering the proximity to Argentina (R genotype). The importation of European queens from the United States, Europe and Argentina by Brazilian beekeepers and proximal geographical contact with Argentina (natural swarm dispersion) itself could have been two ways for this genotype to arrive in Brazil.

MATERIAL AND METHODS

Polymorphic RAPD-type DNA markers were used to determine the genotypes of V. destructor mites originating from different localities in Brazil and from other American countries.

Samples of V. destructor from 5 States of Brazil were analyzed: Santa Catarina (N = 20), São Paulo (N = 20), Minas Gerais (N = 10), Rio Grande do Norte (N = 10), Amazonas (N = 8), and Fernando de Noronha Island - PE (N = 10). Other varroa mites from American countries were also sampled: Argentina (N = 20), Venezuela (N = 10), Cuba (N = 7), Chile (N = 10), Uruguay (N = 15), Colombia (N = 5), and Mexico (N = 15).

Female adult mites were used for DNA analysis. The mites were collected from worker and/or drone broods and immediately transferred to 70% alcohol. In the laboratory, the mites were removed from the alcohol, dried and stored in a freezer at -20°C.

Total DNA was extracted individually from each female of V. destructor using the fast method adapted from Anderson and Fuchs (1998). After washing in 70% ethanol, the mite was transferred to a microcentrifuge tube containing 40 µL 2X lysis buffer (120 µg/mL proteinase K, 0.1 M KCl, 0.02 M Tris-HCl, pH 8, 3.5 mM MgCl$_2$, 0.9% Tween 20, 0.9% NP40 and 0.02% gelatin), which was incubated first at 65°C for 30 min, and 95-100°C for 10 min, and finally, the content was diluted with dH$_2$O.

The primer OPE-07 (5'AGA TGC AGC C 3') (Kraus and Hunt, 1995) was used for DNA amplification and to detect the R and J genotypes.

The samples were prepared for amplification in a volume of 10 µL (Kraus and Hunt, 1995; modified). The reaction mix contained 10 mM Tris-HCl, 50 mM KCl, 2 mM MgCl$_2$, 0.1 mM of each dNTP, 0.2 µM of the primer, 0.5 U Taq polymerase, and 1.5 µL of the extracted DNA. The amplification reaction consisted of 48 cycles of 1 min at 94°C, 1 min at 35°C, and 2 min at 72°C. The products were submitted to 1% agarose gel electrophoresis and stained with ethidium bromide (Maniatis et al., 1982). The gels were visualized and photographed under UV light (Doc-Print, Sony and Biosystem Apparatus), and the amplification products were identified using the φX174 RF DNA/HaeIII marker (Invitrogen).

RESULTS AND DISCUSSION

Figure 1 shows the different genotype patterns determined in this study. All samples from Fernando de Noronha Island (Pernambuco State, Brazil) showed the J pattern. Analysis of mites from the southern region of Brazil (Santa Catarina) and other regions (Amazonas, São Paulo, Minas Gerais, and Rio Grande do Norte) showed the R pattern. Mites from other American countries (Argentina, Uruguay, Colombia, Cuba, Chile, and Mexico) also showed only the R genotype.
The results confirmed the initial hypothesis of the study that the R genotype was already present not only in the south but also in the southeast, northeast (except Fernando de Noronha Island) and north region of Brazil as well as other countries examined, with mites showing the same genetic pattern (RAPD) as that found in Argentina, Canada and United States (de Guzman and Rinderer, 1998).

As there is no barrier to the dispersion of the varroa in all the continental part of Brazil, the increase in the reproductive ability of the mite corroborates with the hypothesis of genotype substitution occurring over time. This hypothesis is reinforced by the presence of the J genotype on Fernando de Noronha Island. The Italian bees were introduced on this island in 1984, when some colonies of bees with imported Italian queens (*Apis mellifera ligustica*) of the United States had been taken from Ribeirão Preto, State of São Paulo, to this island (Malgodi et al., 1986). Therefore, the geographic isolation must explain the exclusive occurrence of the J genotype on Fernando de Noronha Island.

Mites continue to expand throughout the world and the constant changes in the world (collective apiculture) may lead to new crossing opportunities for mites of different origins. New combinations of genetic markers provide evidence for this hybridization, as already observed in some regions of the United States and Canada with the occurrence of the two genotypes R and J in the same bee colony (de Guzman and Rinderer, 1999). This event was also examined in the present study, and our results do not indicate the presence of the two *V. destructor* genotypes in the same Africanized bee colony.

Using mitochondrial markers, Garrido et al. (2003) determined the relationship between fertility and haplotype in a study of *Varroa destructor* mites sampled from colonies of *A. mellifera carnica* and Africanized honey bees (*Apis mellifera*) in Germany and Brazil, respectively. Both in Germany and in Brazil, only the *V. destructor* Korea haplotype was found, though the Japan-Thailand haplotype was previously thought to have been more abundant in Brazil (Anderson, 2000; Anderson and Trueman, 2000).

The fertility of *Varroa* mites in Brazil has increased to European levels. In 1986-1987, only 35% of the varroa females that invaded worker broods had left at the least one viable...
descendant, compared to 72% in 2005-2006 (Carneiro et al., 2007).

It is unknown whether the severity of the effects caused by the Varroa parasite depends on the genotype of the bees and/or on the genotype of the mite. Evidence suggests that the mite V. destructor is a complex species (Kraus and Hunt, 1995; de Guzman et al., 1997; Anderson and Fuchs, 1998; Anderson, 2000; Anderson and Trueman, 2000).

Studies have shown a correlation between the various genotypes of the mite and its fertility in different geographical regions (Garrido et al., 2003). This information may be helpful in understanding the diverse effects and relationships of the mite with bees in different regions of the world. DNA analysis by the RAPD and mtDNA techniques has permitted a precise study of genetic variability among mite populations.

However, the temporal changes in mite fertility and genotype (RAPD), haplotype (mtDNA) that occur in Brazil are not fully correlated with the increasing infestation levels observed at the moment. Other mechanisms limiting mite populations include removal of brood infested with Varroa (hygienic behavior) (Guerra Jr. et al., 2000), mite mortality in adult bees (Correa-Marques et al., 2002; Moretto, 2002) and mortality of mite offspring (Mondragón et al., 2006). These factors among others determine the tolerance of these bees to the mite Varroa destructor and keep the levels of infestation in Brazil low. It is not known if the substitution and/or genetic alteration of genotype (RAPD) and haplotype (mtDNA) is recent. Although the infestation levels of Varroa destructor in Brazil are low, a monitoring program is very important.

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