Genetic parameters for five traits in Africanized honeybees using Bayesian inference

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Abstract

Heritability and genetic correlations for honey (HP) and propolis production (PP), hygienic behavior (HB), syrup-collection rate (SCR) and percentage of mites on adult bees (PMAB) of a population of Africanized honeybees were estimated. Data from 110 queen bees over three generations were evaluated. Single and multi-trait models were analyzed by Bayesian Inference using MTGSAM. The localization of the hive was significant for SCR and HB and highly significant for PP. Season-year was highly significant only for SCR. The number of frames with bees was significant for HP and PP, including SCR. The heritability estimates were 0.16 for HP, 0.23 for SCR, 0.52 for HB, 0.66 for PP, and 0.13 for PMAB. The genetic correlations were positive among productive traits (PP, HP and SCR) and negative between productive traits and HB, except between PP and HB. Genetic correlations between PMAB and other traits, in general, were negative, except with PP. The study permitted to identify honeybees for improved propolis and honey production. Hygienic behavior may be improved as a consequence of selecting for improved propolis production. The rate of syrup consumption and propolis production may be included in a selection index to enhance honeybee traits.

Keywords: heritability, genetic correlations, environmental effects, Apis mellifera.

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Introduction

Apiculture in Brazil is based on the well adapted and productive Africanized honeybee (Silva and Freitas, 2004). The world leaders in honey production are China, USA, Argentina and Mexico (Bölke and Palmeira, 2006). Brazil, according to same authors, is the eleventh largest honey producer and the fifth exporter in worldwide, but has potential to increase both production and productivity. According to Aizen and Harder (2009), the global population of managed honeybee hives has increased ~45% during the last 50 years, but no data are available on feral populations. These authors state that increasing demands for agricultural pollination services are driving the increase in honeybee populations, but there has been a much more rapid (> 300%) increase in the fraction of agriculture that depends on animal pollination in the same period, stressing a need for global pollination capacity. This implies a need to increase honeybee numbers, survival and productivity, and this can be achieved through applying animal breeding theory to managed populations. This situation also exists in Brazil (Guedes et al., 2011). However, information on environmental and genetic factors that affect economic traits in the different genotypes of honeybees across the country is limited.

The Africanized honeybee is considered more resistant to diseases and more tolerant to Varroa destructor than the European honeybees, breeding for improvement in productive and resistance/tolerance traits should have significant economic impact on this activity (Büchler et al., 2010). Honey production, production of propolis, foraging behavior, hygienic behavior and lower percentages of mites on adult bees (or grooming behavior) should be included in genetic evaluations and selection programs in order to choose the best queens to increase honey or propolis production and productivity (Milne Jr, 1985; Bienefeld and Pirchner, 1990; Thimann and Manrique, 2002; Stanimirovic et al., 2010; Wilson-Rich et al., 2010). However, before selection can be carried out, it is necessary to estimate the heritabilities and the genetic correlations between productive and...
resistance/tolerance traits to predict the response to selection for a given trait.

There are few reports on parameter estimates in Africanized honeybees in Brazil (Faquinello et al., 2011; Costa-Maia et al., 2011). Therefore, the objective of this study was to estimate the heritabilities and genetic correlations for honey production, propolis production, syrup collection rate, hygienic behavior and percentage of mites on adult bees in a population of Africanized honeybees in southern Brazil.

Material and Methods

The study was conducted from April 2009 to October 2010 at the Agronomic Experimental Station of the Federal University of Rio Grande do Sul (EEA/UFRGS), in Eldorado do Sul, Rio Grande do Sul, Brazil. Africanized honey bee queens used in this study were donated by a local beekeepers association. A total of 110 queens were evaluated over three generations (40, 40 and 30 queens per generation). The queens were raised from each of the first 40 colonies for the second generation, and from each of the second generation colonies for the third one. Missing records in the third generation were due to problems in an incubator for queen rearing and loss of some colonies over winter because of falling trees in an apiary. Queen rearing was realized by the larval transference method. The new queens were marked with enamel paint and introduced into the colonies two days after the laying queen had been removed. The new queens were mated naturally. The first measurement was carried out at least 65 days after the new queens started to lay eggs.

Honey production was measured twice, in May 2009 and March 2010. To measure honey production, frames were weighed before and after centrifugation. Syrup collection rate, used as a proxy for estimating hoarding behavior, was measured in July/August 2009, February and August 2010, totalling three measurements. All the hives had the same numbers of empty combs, but syrup storage was not measured, because the objective was to determine the rate of consumption. Glass recipients (600 mL) covered by metal lids with eight holes (1.5 mm width) were used. Each recipient was completely filled with water:sugar:honey syrup in a 4:4:2 proportion. Two (July/August) to three (on February) recipients were introduced into the hive in the afternoon. After 12 hours, leftover syrup was measured to determine the rate of consumption.

Measurements of propolis production were carried out in July and August 2009 (winter), April and May 2010 (autumn) and September/October 2010 (spring). Production was measured using propolis collectors on two sides of the hive (22 cm x 3 cm). These were covered with a transparent adhesive tape. This tape was used to protect honeybees from environmental conditions and predators. The collectors remained in the hives for 25 days before the propolis was cut and weighed.

Measurement of hygienic behaviour was realized by the Needle Test (Olszewski and Paleolog, 2007). On a comb containing pupae in the initial phase of eye pigmentation or light pink colour (10 to 14 days age) 100 cells were pricked simultaneously with a “brush” containing 100 needles. In cases of irregularly constructed combs, the cells were pricked individually with a single needle. A similar nearby area (not pricked) of the comb was used to correct for natural cleaning of parasitized or sick pupae. The comb was photographed before the test and 24 hours thereafter to count the number of perforated cells and (un)cleaned combs. To guarantee the quality of images, the same comb was photographed in the field four times to evidence the difference between eventual loads of pollen and perforated pupae that were not cleaned by the bees. Thus hygienic behaviour was evaluated as:

\[ HB = \left( \frac{EC_1 - EC_2}{OC} \right) \times 100 - \left( \frac{A - B \times 100}{A} \right), \]

where A is the number of cells with operculated pupae (witness area; not pricked); B is the number of empty cells of the witness area before the needle test; C is the number of empty cells of witness area after the needle test; \( EC_1 \) is the number of empty cells in the test area 24 h after the needle test; \( EC_2 \) is the number of empty cells in test area before the needle test; \( OC \) is the number of operculated cells (with pupae) in the test area before the needle test.

The measurement of percentage of mites on adult bees was adapted from De Jong et al. (1982). A solution of a neutral commercial detergent in water was prepared (0.5 mL/L). Recipients (250 mL) were filled with 100 mL of this solution and taken to the field. Samples of adult honeybees from the brood area of the combs were picked at the same time as the hygienic behaviour test. The samples were shaken for 1 min in a glass recipient (600 mL) and sieved to separate mites and honeybees.

The quantity of honeybees in a hive may influence the performance of the colony (Topal et al., 2008) and is referenced in literature as colony strength (Nasr et al., 1990). The strength of the colony was measured by counting the number of frames covered with adult bees. This measurement was taken for all traits to be used as covariate in the statistical models. Data were analyzed to verify normality using the Univariate Procedure of the SAS program (SAS Institute Inc, Cary, North Carolina). Phenotypic records were transformed (Square root for honey production, cubic root for propolis production and syrup collection rate, and logarithm of percentage of mite + 1). The GLM Procedure was used to check the influence of non-genetic effects on traits, and which ones were to be considered in the models to estimate genetic parameters. The description of the data set used in the study is shown in Table 1.

The Sire-Maternal Grandsire Model (MGS) was used to estimate genetic parameters, considering the relationship
among queens. Models included non-genetic effects of location of the hive, season-year, and number of frames covered with bees for honey production, syrup collection rate, and percentage of mites on adult bees. For propolis production and hygienic behaviour the localization of the hive and the number of frames covered with bees were considered. The genetic and residual effects were considered for all traits.

The matricial model used to estimate genetic parameters in a single-trait analysis was:

\[ y = X\mathbf{\beta} + \mathbf{Z}_a + \mathbf{e}, \]

where \( y \) is the vector of observations; \( \mathbf{\beta} \) is the vector of non-genetic effects; \( \mathbf{a} \) is the vector of animal effects; \( \mathbf{e} \) is the vector of residual effects; \( X \) is the design matrix of order \( n \times p \), which relates records to non-genetic effects; \( \mathbf{Z}_a \) is the design matrix of the order \( n \times q \), which relates records to animal effects. The matricial model used to estimate genetic parameters in multi-trait analysis was:

\[
\begin{bmatrix}
\mathbf{y}_1 \\
\mathbf{y}_2
\end{bmatrix} =
\begin{bmatrix}
X_1 & 0 \\
0 & X_2
\end{bmatrix}
\begin{bmatrix}
\mathbf{b}_1 \\
\mathbf{b}_2
\end{bmatrix} +
\begin{bmatrix}
\mathbf{Z}_1 & 0 \\
0 & \mathbf{Z}_2
\end{bmatrix}
\begin{bmatrix}
\mathbf{u}_1 \\
\mathbf{u}_2
\end{bmatrix} +
\begin{bmatrix}
\mathbf{e}_1 \\
\mathbf{e}_2
\end{bmatrix},
\]

where \( \mathbf{y}_1 \) is the vector of observations to the \( i \)-th trait; \( \mathbf{b}_1 \) is the vector of non-genetic effects to the \( i \)-th trait; \( \mathbf{a} \) is the vector of animal effects to the \( i \)-th trait; \( \mathbf{e}_1 \) is the vector of residual effect of the \( i \)-th trait; \( X_1 \) and \( \mathbf{Z}_1 \) are incidence matrices that relate the observations of the \( i \)-th trait to environmental and animal effects. It was assumed that:

\[
\begin{bmatrix}
\mathbf{a}_1 \\
\mathbf{a}_2 \\
\mathbf{e}_1 \\
\mathbf{e}_2
\end{bmatrix} =
\begin{bmatrix}
g_{11}A & g_{12}A & 0 & 0 \\
g_{21}A & g_{22}A & 0 & 0 \\
0 & 0 & r_{11} & r_{12} \\
0 & 0 & r_{21} & r_{22}
\end{bmatrix},
\]

where \( g_{ij} \) is the additive genetic variance to direct effects of the trait \( i \); \( g_{ij} \) are the additive genetic covariance between traits; \( g_{22} \) is the additive genetic variance to direct effects of the trait \( 2 \); \( r_{ij} \) are the elements of \( \mathbf{R} \), the (co)variance matrix to residual effects.

Components of variance and genetic parameters were estimated using Bayesian Inference by MTGSAM software (Multiple Trait Gibbs Sampling in Animal Models) developed by Van Tassell and Van Vleck (1995). The initial values for the components of variance were obtained using REML analysis and values published in the literature (Rothenbuhler et al., 1979; Bienefeld and Pirchner, 1990; Stanimirovic et al., 2008). In the MTGSAM analyses, the initial distribution was assumed as symmetric and the definition parameter of the initial distribution was assumed as 4 (single-trait analysis) and 5 (multi-trait analysis).

The models were analyzed, first by generating two "pilot-samples", which were monitored by the Raftery and Lewis (1992) diagnostic test to verify convergence. The test, implemented in the BOA software package (Bayesian Output Analysis) generated indications of burn-in, number of iterations and thinning interval. These indications were used to generate 20,000,000 (single-trait) and 100,000,000 (multi-trait) total iterations, a burn-in of 3,000,000 (single and multi-trait analysis) and thinning interval of 1,000 (single-trait) and 2,000 (multi-trait) iterations, which resulted in 48,500 (single-trait) and 17,000 (multi-trait) final samples. The convergence criterion used was 10^-12. Convergence of samples was monitored using values of serial correlation by graphical analysis and diagnostic tests of Raftery and Lewis (1992), Geweke (1992) and Heidelberger and Welch (1983), and values of Markov Chain Error, calculated by the package BOA (Bayesian Output Analysis) (Smith, 2005), implemented in R software (R Development Core Team, 2010). The highest probability density interval for estimation of components of variance and genetic parameters was tested.

**Results**

The local effect (Table 2) influenced syrup collection rate, propolis production and hygienic behavior (\( p < 0.05 \)). The season-year effect was significant (\( p < 0.001 \)) only for syrup-collection rate. The number of frames with adult bees, fitted as a covariate, was significant (\( p < 0.05 \)) for honey production, syrup collection rate and propolis production. The effects in the model explained 24%, 32%, 41%, 66% and 74% of the total variation for honey production (HP), percentage of mites on adult bees (PMAB), hy-

**Table 1** - Description of the data set used in the analysis.

| Traits                              | Mean ± sd\(^1\) | CV % | Shapiro-Wilk test | Mean ± sd\(^2\) | CV % | Shapiro-Wilk test |
|-------------------------------------|------------------|------|-------------------|------------------|------|-------------------|
| Honey production                    | 10.72 ± 6.63 kg  | 61.88| \( p < 0.001 \)  | 3.11 ± 1.02      | 32.88| \( p = 0.47 \)   |
| Syrup collection rate               | 49.42 ± 35.22 mL/h | 71.26| \( p < 0.001 \)  | 8.14 ± 1.93      | 23.73| \( p = 0.03 \)   |
| Propolis production                 | 32.97 ± 32.95 g  | 99.94| \( p < 0.01 \)   | 2.79 ± 1.01      | 36.46| \( p = 0.12 \)   |
| Hygienic behaviour                 | 69.90 ± 22.92%   | 32.78| \( p = 0.0003 \) | -                | -    | -                 |
| Percentage of mites on adult bees   | 3.86 ± 2.96%     | 76.68| \( p < 0.01 \)   | 1.44 ± 0.58      | 40.00| \( p = 0.46 \)   |
| Number of frames with adult bees   | 8.06 ± 3.73      | 46.28| \( p < 0.001 \)  | 1.95 ± 0.31      | 15.61| \( p = 0.002 \)  |

\(^1\)original data, \(^2\)transformed data, CV% = coefficient of variation, sd = standard deviation.
gienic behavior (HB), propolis production (PP) and syrup collection rate (SCR), respectively.

Heritability estimates (Table 3) were 0.16 for HP, 0.23 for SCR, 0.66 for PP, 0.52 for HB and 0.13 for PMAB on non-transformed data, and 0.17 for HP, 0.29 for SCR, 0.67 for PP and 0.14 for PMAB for transformed data (Table 4). The transformation of data did not influence the results for the mean heritability and highest probability density intervals (HPDI). Thus, non-transformed data were used to estimate heritability and genetic correlations. In general, HPDI showed large intervals for variance components and heritability. These results may be due to the low number of queens evaluated and/or to the natural mating scheme used.

The means of the posterior distributions of the genetic correlations (Table 5) were negative between HP/SCR, HB and PMAB. On the other hand, the residual correlations between these same traits were positive. The genetic correlations between HP, SCR and PP, and between PP with HB and PMAB were positive. In general, residual correlations

| Table 2 - Mean squares and significance levels for traits in Africanized honey bees. |
|---------------------------------|------------|----------------|------------|----------------|----------------|----------------|
|                                 | df HP      | df SCR         | df PP      | df HB          | df PMAB        |               |
| Local                           | 4 0.97 ns  | 4 8.79 *       | 4 3.35 **  | 4 1636 *       | 4 0.33 ns      |               |
| Season-year                     | 1 1.12 ns  | 3 69.37 **     | 2 0.74 ns  | 3 1217 ns      | 2 0.56 ns      |               |
| No. of frames                   | 7 2.06 *   | 18 3.90 **     | 18 1.10 *  | 17 578 ns      | 18 0.38 ns     |               |
| Error                           | 74 68.45   | 88 111.38      | 52 26.56   | 57 28.26       | 86 24.90       |               |
| R²                              | 0.24       | 0.74           | 0.66       | 0.41           | 0.32           |               |

* = significant (p < 0.05), ** = highly significant (p < 0.001), ns = non-significant, df = degree of freedom, R² = coefficient of determination, HP = honey production, PMAB = percentage of mites on adult bees, HB = hygienic behavior, PP = propolis production, SCR = syrup-collection rate.

| Table 3 - Posterior distributions of the variance components for five traits in Africanized honey bees (not transformed) with highest probability density. |
|---------------------------------|------------|----------------|------------|----------------|----------------|
|                                 | HP         | SCR            | PP         | HB             | PMAB           |
| σ²_x                           | 6.53 (0.58; 20.48) | 1.58 (0.19; 4.32) | 470.55 (118.86; 773.35) | 303.06 (46.92; 572.81) | 0.99 (0.08; 3.38) |
| σ²_e                           | 33.34 (15.41; 50.21) | 5.33 (2.23; 7.9) | 234.36 (35.05; 546.28) | 276.13 (43.19; 536.06) | 6.65 (3.43; 9.45) |
| σ²_p                           | 40.27 (28.9; 52.65) | 6.92 (5.31; 8.76) | 704.91 (500.22; 933.09) | 579.19 (411.1; 768.44) | 7.64 (5.78; 9.73) |
| h²                             | 0.16 (0.02; 0.52) | 0.23 (0.03; 0.61) | 0.66 (0.24; 0.94) | 0.52 (0.12; 0.89) | 0.13 (0.01; 0.45) |

HP = honey production, SCR = syrup collection rate, PP = propolis production, HB = hygienic behavior, PMAB = percentage of mites on adult bees.

| Table 4 - Posterior distributions of the variance components for four traits in Africanized honey bees (transformed) with highest probability density intervals. |
|---------------------------------|------------|----------------|------------|----------------|----------------|
|                                 | HP         | SCR            | PP         | PMAB           |
| σ²_x                           | 0.17 (0.02; 0.52) | 0.47 (0.06; 1.17) | 0.39 (0.09; 0.64) | 0.04 (0.004; 0.12) |
| σ²_e                           | 0.80 (0.35; 1.19) | 1.15 (0.40; 1.78) | 0.19 (0.03; 0.45) | 0.24 (0.13; 0.33) |
| σ²_p                           | 0.96 (0.70; 1.27) | 1.62 (1.23; 2.04) | 0.59 (0.42; 0.78) | 0.28 (0.21; 0.35) |
| h²                             | 0.17 (0.002; 0.54) | 0.29 (0.04; 0.69) | 0.67 (0.25; 0.94) | 0.14 (0.02; 0.42) |

HP = honey production, SCR = syrup collection rate, PP = propolis production, HB = hygienic behavior, PMAB = percentage of mites on adult bees.

| Table 5 - Means of posterior distributions of the genetic (superior diagonal) and residual correlations (inferior diagonal) with the highest probability density intervals. |
|---------------------------------|------------|----------------|------------|----------------|----------------|
| HP                              | SCR        | PP             | HB         | PMAB           |
| HP                              | -          | 0.48 (-0.35; 0.96) | 0.58 (-0.04; 0.94) | -0.34 (-0.09; 0.47) | -0.52 (-0.98; 0.63) |
| SCR                             | -0.13 (-0.75; 0.35) | - | 0.67 (0.23; 0.94) | -0.24 (-0.8; 0.45) | -0.66 (-0.98; 0.04) |
| PP                              | -0.44 (-0.87; 0.09) | -0.09 (-0.70; 0.92) | - | 0.15 (-0.41; 0.63) | 0.49 (-0.15; 0.91) |
| HB                              | 0.36 (-0.30; 0.87) | 0.80 (0.45; 0.97) | 0.65 (0.03; 0.97) | - | -0.82 (-0.98; -0.44) |
| PMAB                            | 0.09 (-0.49; 0.57) | 0.30 (-0.07; 0.76) | -0.59 (-0.92; -0.14) | 0.30 (-0.30; 0.85) | - |

HP = honey production, SCR = syrup collection rate, PP = propolis production, HB = hygienic behavior, PMAB = percentage of mites on adult bees.
were negative, except between PP and PMAB. The genetic correlations between PP and SCR and between HB and PMAB were more precisely estimated, showing the smallest intervals in HPDI. The same precision was observed in residual correlations between HB and SCR, between HB and PP and between PP and PMAB.

**Discussion**

As there is little information on environmental and genetic parameters for Africanized honeybees in the literature we compared our results with studies on European honeybees, although these used different methods and number of queens or records. Bar-Cohen et al. (1978) evaluated 562 colonies of European honeybees, using regression methods. Bienefeld and Pirchner (1990) evaluated 5581 records of European bees, using the REML method. In Brazil, Costa-Maia et al. (2011) and Faquinello et al. (2011) used Bayesian inference to evaluate 40 and 193 Africanized honeybees, respectively.

The location of the hive influenced PP, HB and SCR and the season-year was an important effect for SCR. These effects are related to the distance between the hive and the source of pollen or nectar (Sande et al., 2009), as well as meteorological factors (rain, temperature, clouds) (Holmes, 2002). Although the location was not significant for all traits in this study, it was used in all models, because analyses including this effect showed higher precision in the estimates of heritability and HPDI. Year-season was included in models for HP and PMAB. The number of frames covered with bees influenced PP, HP and SCR, which are traits related to production. Topal et al. (2008) reported that the number of frames with bees had a significant positive relationship with honey yield. Thus, this effect was included in all models as linear covariate.

The heritability (0.16) for HP was lower than the value reported by Bar-Cohen et al. (1978) for Italian bees (0.54), but higher than the low value found by Praagh et al. (2006) for Carnolian honeybees (0.03). While these parameters were obtained by regression methods, Bienefeld and Pirchner (1990) used the Restricted Maximum Likelihood method (REML) for *Apis mellifera carnica* and found 0.15 and 0.26 for effects of queen and worker bees, respectively. These differences among parameter estimates may be due to different strains or races of honeybees, different methods of estimation and/or environmental effects considered in the models. Honey production is one of many economically valuable traits, being greatly influenced by environmental factors such as a regular supply of pollen and nectar over the season, a hive fully populated with worker bees, the weather, and the quality of care that honeybees receive from beekeepers (Holmes, 2000). The low heritability found in this study may be explained by the residual variance, which was five times higher than the genetic variance (Table 3).

The heritability (0.23) for SCR was lower than that reported for rapid (0.55) and slow (0.32) lines selected for syrup collection under laboratory conditions by Rothenbuhler et al. (1979) and higher than the heritability (0.187 ± 0.029) found by Milne Jr (1985) for the removal of sugar syrup from vials and the storage in combs by caged honey bees. Despite the methodological difference between the field test in the present study and field tests using flight cages or laboratory tests (Olszewski and Paleolog, 2005), the result (heritability) appears to be quite similar, this indicating that it is possible to obtain genetic progress using this kind of hoarding behavior measure as a selection criterion.

Some studies have shown that selection for improving propolis production in Africanized bees is feasible (Manrique and Soares, 2002; Thimann and Manrique, 2002). However, numerous studies have focused mainly on the chemical components of propolis that could be used to treat human diseases (Simone-Fingston and Spivak, 2010), and others on the importance of propolis to bee health (Simone et al., 2009). The heritability (0.66) estimated for PP is high and should permit genetic improvement through selection.

The heritability estimate (0.52) for HB was lower than the value (0.63) found for Carnolian honeybees by Stanimirovic et al. (2008). Costa-Maia et al. (2011), using Bayesian inference, estimated a lower heritability value (0.28) for Africanized bees in Brazil. Hygienic behavior is widely studied and documented in the literature (Wilson-Rich et al., 2009). There are several breeding programs and/or institutions involved in the selection for more hygienic queens/worker bees to obtain more healthy and productive colonies (Stanimirovic et al., 2008; Bourgeois and Rinderer, 2009; Costa-Maia et al., 2011).

The low heritability estimated for PMAB (0.13) is explained by the low genetic variance, which was 6.7 times lower than the residual variance. Despite the slow genetic progress that may be achieved, the selection of honeybees with low levels of mites or with tolerance to *Varroa destructor* is economically very important (Spivak and Reuter, 1998; Martin and Medina, 2004; Rinderer et al., 2010). While this mite has killed millions of colonies of European honeybees in Europe and North America, the Africanized bee has a certain tolerance to this mite (Martin and Medina, 2004). Nevertheless, the presence of the mite may decrease the productivity of the colony.

The genetic correlations between HP, SCR and PP (0.48, 0.58, and 0.67) were positive and favorable, which suggests that the selection for SCR or PP will increase HP. Certain studies in the literature have shown a positive phenotypic correlation between the amounts of syrup gathered or collected by bees (hoarding behavior) in the laboratory/field and the honey production in the hives (Olszewski and Paleolog, 2005; Zárate et al., 2008). Another study also found a positive correlation (Pearson) between PP and HP.
in Africanized honeybees, with $r = 0.422$ ($p = 0.00001256$), showing that the bees that produced more propolis also produced more honey (Manrique and Soares, 2002). The negative residual correlations among the same traits (-0.13, -0.44, -0.09) indicate that the improvement of environmental conditions will have an antagonistic effect.

Genetic correlations between HP and HB (-0.34) and between SCR and HB (-0.24) were antagonistic, which suggest a negative response in selection. Furthermore, the residual correlations (0.36, 0.8) were positive, which indicates that environmental conditions WERE shared. These results are different from those reported by Spivak and Reuter (1998). These authors compared naturally mated queens from a hygienic line of Italian honey bees (Apis mellifera ligustica) with colonies of a commercial line of Italian bees not selected for hygienic behavior. They found that the hygienic colonies removed more freeze-killed brood (hygienic behavior) and produced significantly more honey than the commercial lines.

The positive genetic correlation between PP and HB (0.15) differs from other estimates in this study and was not expected. Although the two traits represent different trends, and at least apparently the genetic association is not clear or not documented in the literature, these results suggest that the selection for more hygienic bees may increase propolis production. Moreover, the positive residual correlation (0.65) was high and suggests that environmental effects affect both traits.

The genetic correlations between PMAB with HP, SCR and HB (-0.52, -0.66, -0.82) were high and favorable, except between PMAB and PP (0.49), which was in opposite direction. The results indicate that selection for HB, HP/SCR should decrease PMAB, but the selection for PP should increase PMAB. Many studies have shown the same relationship between HB, the tolerance to Varroa destructor, and HP (Spivak and Reuter, 1998; Harris, 2008). These same studies reported that HB in general and HB towards Varroa destructor (Varroa-Sensitive Hygienic Behavior) are mechanisms of tolerance against diseases and varroa mites. Spivak and Reuter (1998) reported that hygienic bees (genetically selected) also had significantly less chnkbrood, no American foulbrood, fewer mites, and produced more honey than commercial (not selected) bees. Basically, there are two mechanisms that affect the levels of infestation of mites: grooming behavior (adult bees detect and remove mites from themselves and from nestmates) and HB (the detection and removal of dead/ill or infested pupae from capped brood cells by adult bees). Residual correlations (0.09, 0.30 and 0.30) between PMAB with HP, SCR and HB were in opposite direction and between PP and PMAB (-0.59) they were in the same direction.

In conclusion, the studied population of Africanized honeybees showed sufficient genetic variation to begin a breeding program. Depending on the breeding approach in a future program, commercial stocks of Africanized honeybees that produce more propolis and honey may be achieved. Moreover, hygienic behavior may be improved as a consequence of selecting for improved propolis production, but additional studies should be realized to better understand the genetic association between these traits. On the other hand, the rate of syrup consumption and propolis production may be used as selection criteria, preferably by including these traits in a selection index to improve traits of honeybees in breeding programs.

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