Relevant genetic differentiation among Brazilian populations of *Anastrepha fraterculus* (Diptera, Tephritidae)

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Abstract

We used a population genetic approach to detect the presence of genetic diversity among six populations of *A. fraterculus* across Brazil. To this aim, we used Simple Sequence Repeat (SSR) markers, which may capture the presence of differentiative processes across the genome in distinct populations. Spatial analyses of molecular variance were used to identify groups of populations that are both genetically and geographically homogeneous while also being maximally differentiated from each other. The spatial analysis of genetic diversity indicates that the levels of diversity among the six populations vary significantly on an eco-geographical basis. Particularly, altitude seems to represent a differentiating adaptation, as the main genetic differentiation is detected between the two populations present at higher altitudes and the other four populations at sea level. The data, together with the outcomes from different cluster analyses, identify a genetic diversity pattern that overlaps with the distribution of the known morphotypes in the Brazilian area.

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Keywords
Anastrepha fraterculus, microsatellites, population genetic differentiation, morphotypes

Introduction

The South American fruit fly *Anastrepha fraterculus* Wiedemann (Diptera: Tephritidae) belongs to the *fraterculus* group, which comprises a total of 34 formally described species (Norrbom et al. 2012) that can be distinguished only by minor morphological characters. The major diagnostic character for species within this group is the aculeus apex, which shows a large degree of intraspecific variation (Araujo and Zucchi 2006) as a result of genetic and environmental factors (Aluja 1994, Smith-Caldas et al. 2001).

The nominal species *A. fraterculus* is widely distributed from the Rio Grande Valley in northern Mexico to central Argentina, infesting over 100 hosts (Norrbom 2004), thus being a species of major economic importance in Brazil and other countries in South America (Steck 1991, Steck 1999, Zucchi 2008). Its distribution in South America was thought to be in two broad and unconnected bands along the east coast and the western and northern edges of the continent with a hiatus comprising the Amazon basin (Steck 1999, Malavasi et al. 2000). Yet, recent collections have recorded this species in the Brazilian Amazon in the states of Amapá, Pará, Tocantins, and Maranhão infesting 10 different hosts (Zucchi et al. 2011).

*A. fraterculus* has long been reported to show extensive morphological variation along its geographic distribution (Lima 1934, Stone 1942, Steck 1991, 1999). In his taxonomic review of the genus *Anastrepha*, Stone (1942) stated “it is possible that it will eventually be found to represent a complex of species rather than a single one”. Since then, a good deal of research has documented and concluded that the nominal species *A. fraterculus* actually comprises an unresolved complex of cryptic species. Evidence comes from studies on morphological variation (Steck 1999, Hernández-Ortíz et al. 2012, for review), multivariate morphometric analyses (Hernández-Ortíz et al. 2004, 2012), differences in host use (Steck 1999, for review), behaviour (Yamada and Selivon 2001, Selivon et al. 2005, Vera et al. 2006, Vaničková et al. 2015), the presence and degree of reproductive isolation (Devescovi et al. 2014 and references therein), and genetic analyses (Silva and Barr 2008, for review). However, the actual number of species within the *A. fraterculus* complex and their distribution is yet to be elucidated.

Genetic studies performed on *A. fraterculus* populations so far have revealed the following putative biological entities based on geography: an Andean lineage (Steck 1991), a Mexican species (Steck 1991, Smith-Caldas et al. 2001, Barr et al. 2005 and the morphometric studies of Hernández-Ortíz et al. 2004, 2012), a Guatemalan lineage (Smith-Caldas et al. 2001), a second Venezuelan species (Smith-Caldas et al. 2001), a Peruvian lineage (Steck 1991), and three Brazilian species (Morgante et al. 1980, Smith-Caldas et al. 2001, Selivon et al. 2005, and Ruiz et al. 2007a, 2007b) (Silva and Barr 2008, for review). In addition, morphometric studies revealed that Mexican populations formed a single morphotype, which was distinct from South
American populations (Hernández-Ortíz et al. 2004). Hernández-Ortíz et al. (2004) also identified an Andean morphotype (one population from Colombia) and a Brazilian morphotype comprising two populations from Brazil (states of São Paulo and Santa Catarina) and a population from Argentina (Tucumán). A study using isozymes, karyotypes, morphometry, and crossings on populations of *A. fraterculus* from Brazil recognized two species, *Anastrepha* sp.1 aff. *fraterculus* and *Anastrepha* sp.2 aff. *fraterculus* (Selivon et al. 2005). More recently, a multivariate morphometric analysis comprising 32 *A. fraterculus* populations identified seven distinct morphotypes: a Mexican morphotype, a Venezuelan morphotype, an Andean morphotype, a Peruvian morphotype, and three Brazilian morphotypes (Brazilian 1, Brazilian 2, and Brazilian 3) (Hernández-Ortíz et al. 2012). Although Brazilian populations of the nominal species *A. fraterculus* most likely comprise at least three morphotypes, published studies are based mostly on samples from the south-eastern region, while very few populations from other regions of the country have been examined.

Previous genetic studies using DNA sequencing of mitochondrial genes from different populations suggested that both the *fraterculus* group and the *A. fraterculus* complex have a recent evolutionary history, and thus molecular markers with a higher power of resolution were required to help understand the specific/subspecific differentiation within and between populations of this group (McPherson et al. 1999, Smith-Caldas et al. 2001). Microsatellites are a class of highly polymorphic molecular markers widely distributed in the genome of eukaryotes that can be useful to clarify such patterns of gene flow and to identify the spatial locations of genetic discontinuities (population boundaries) in studies of species complexes (Chambers and MacAvoy 2000, Barbará et al. 2007, Aketarawong et al. 2014). Within the Tephritidae, microsatellites have been successfully developed and applied for several *Bactrocera* species (Shearman et al. 2006; Aketarawong et al. 2006, 2007, Augustinos et al., 2008, Virgilio et al. 2010; Drew et al. 2011), for *Rhagoletis cerasi* L. (Augustinos et al. 2011), for a few *Ceratitis* species (Bonizzoni et al. 2001, 2004, Meixner et al. 2002, Baliraine et al. 2003, 2004, Silva et al. 2003, Delatte et al. 2013, Virgilio et al. 2013), for *Anastrepha suspensa* (Loew) (Fritz and Schable 2004, Boykin et al. 2010), and for *Anastrepha obliqua* (Macquart) (Islam et al. 2011). Microsatellites have only recently been isolated in *A. fraterculus* (Lanza-Vecchia et al. 2014) and have proven useful for the analysis of population dynamics and differentiation across the distribution range of this polymorphic species.

From an applied perspective, the correct identification of populations and species is an important step in the implementation of biologically-based control methods such as the Sterile Insect Technique (SIT) against this fruit fly complex (Silva and Barr 2008), which represents a serious constraint for fruit production in South America and a hindrance to the export of fresh fruit from regions where it occurs. Currently, control measures for this pest species rely solely on the use of insecticide cover or bait sprays. Therefore, there is demand for the development of the SIT as it would benefit South American countries such as Argentina, Brazil, and Peru (Cladera et al. 2014). However, the application and efficiency of species-specific control methods such as the SIT are critically dependent on the correct identification of the target pest populations.
and the understanding of the spatial distribution of the pest species, thus the correct
delimitation of species within the *A. fraterculus* complex is paramount.

This paper is centred on the assessment of genetic diversity among *A. fraterculus*
populations from distinct geographic regions across Brazil, most likely belonging to at
least three distinct morphotypes (Silva et al. unpubl. data). For this purpose we used
highly informative SSR markers, which may capture the presence of eventual differenti-
tiative processes across the genome in different populations.

**Methods**

Six populations from three regions across Brazil were sampled from 2007 to 2013 (Table
1, Figure 1). In the Northeastern region, the populations from Monte Alegre (State of
Rio Grande do Norte) and from Una and Porto Seguro (State of Bahia) were sampled.
In the Southeastern region, samples from São Mateus (State of Espírito Santo) and from
Campos do Jordão (State of São Paulo) were examined. In the Southern region, the
population from Vacaria (State of Rio Grande do Sul) was sampled. For each locality,
flies emerging from fruits collected from different trees were considered. Adult females
were identified as *A. fraterculus* by Dr. Elton L. Araujo (UFERSA), Dr. Keiko Uramoto
(USP), Dr. Miguel Francisco Souza Filho (Instituto Biológico) and Dr. Roberto A. Zuc-
chi (USP) using the aculeus shape following Zucchi (2000) (Table 1). Voucher speci-
mens were deposited at the insect collection of the Escola Superior de Agricultura “Luiz
de Queiroz”, USP, Piracicaba, SP, and at the Universidade Estadual de Santa Cruz, Ilhéus,
BA, Brazil. According to the classification of Hernández-Ortíz et al. (2012), the flies col-
lected in Una (BA) can be classified as Brazilian morphotype 3, while those from Vacaria
(RS) and Campos do Jordão (SP) as Brazilian 1 (Silva et al. unpublished data). For the
samples from Monte Alegre, Porto Seguro and São Mateus, no clear-cut information is
available to assign them to a specific morphotype.

**Table 1.** Field collected samples of *Anastrepha fraterculus* Brazilian populations.

| States                  | Sample site       | Morphotype* | Host   | Coordinate          | Elevation |
|------------------------|-------------------|-------------|--------|---------------------|-----------|
| Rio Grande do Norte (RN)| Monte Alegre      | ?           | Guava  | 6.0678W; 35.3322S   | 51.816m   |
| Bahia (BA)             | Una               | 3           | Guava  | 15.2933W; 39.0753S  | 27.737m   |
| Bahia (BA)             | Porto Seguro      | ?           | Guava  | 16.4497W; 39.0647S  | 48.768m   |
| Espírito Santo (ES)    | São Mateus        | ?           | Araçá  | 18.7161W; 39.8589S  | 35.966m   |
| São Paulo (SP)         | Campos do Jordão  | 1           | Raspberry | 22.7394W; 45.5914S | 1627.9m   |
| Rio Grande do Sul (RS) | Vacaria           | 1           | Guava  | 28.5122W; 50.9339S  | 970.79m   |

* The Morphotype classification is based on Hernández-Ortíz et al. (2012)
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Figure 1. Map of the collected samples.
Microsatellite analysis

A total of 171 *A. fraterculus* individuals collected from the above mentioned populations were assessed for their SSR variability. DNA was extracted from three legs of each single fly using the “DNeasy Blood & Tissue” kit (Qiagen, Valencia, CA) following the standard DNeasy protocol. DNA samples were screened using the following ten microsatellite loci: AfD4, AfD105, AfA7, AfA112, AfA115, AfA120, AfA122, AfA117, AfA10, and AfC103 (Lanzavecchia et al. 2014). Allele scoring was performed using an automated ABI PRISM 310 Genetic Analyser (Applied Biosystem) following Aketarawong et al. (2006).

Data analysis

The mean number of alleles (na) and mean null allele frequency (An) (non-amplifying alleles due to changes in the primer binding regions), expected and observed heterozygosity were estimated using GENEPOP version 4.0.7 (Raymond and Rousset 1995) for each population. Deviation from the Hardy-Weinberg equilibrium and linkage disequilibrium, together with their critical levels after the sequential Bonferroni test (Rice 1989), were tested using GENEPOP version 4.0.7 (Raymond and Rousset 1995). The allelic Polymorphic Information Content (PIC) was derived using CERVUS (Kalinowski et al. 2007).

Microsatellite Analyzer (MSA) (Dieringer and Schlötterer 2003) was applied to estimate the pairwise *Fst* values among populations (Weir and Cockerham 1984). The statistical significance of each *Fst* value was assessed by comparing the observed values with the values obtained in 10,000 matrix permutations. Spatial analyses of molecular variance were investigated using SAMOVA 2.0 (Dupanloup et al. 2002). This approach identifies groups of populations that are genetically homogeneous and maximally differentiated from each other without the constraint of being geographically close. The method requires the *a priori* definition of the number of groups (K) of populations that exist and generates F-statistics (*F*sc, *F*st, and *F*ct) using an AMOVA approach. Different numbers of groups (K) were tested, and a simulated annealing procedure permitted the identification of the composition of each of the K groups that maximizes the *F*ct index (proportion of total genetic variance due to differences between groups). The program was run for two to five groups (K = 2 to K = 5) each time with the simulated annealing process repeated 100 times, starting each time with a different partition of the population samples into the K groups. The analysis of molecular variance (AMOVA) was carried out using ARLEQUIN software version 3.11 (Excoffier et al. 2006). Principal Coordinate Analysis (PCoA) in the program GenAIEx 6.5 (Peakall and Smouse 2012) was applied to identify the relationships among populations on the basis of their allele frequencies.
Results

SSR variability

The variability estimates describing the suitability of the ten SSR loci (AfD4, AfD105, AfA7, AfA112, AfA115, AfA120, AfA122, AfA10, and AfC103) for detecting the presence of differentiation among the *A. fraterculus* populations are shown in Table 2. The number of alleles per locus across populations ranges from 7 to 18 with a mean of 12.9, and the mean frequency of null alleles across the loci is generally low (0.04–0.08). Moreover, the Polymorphic Information Content (PIC) estimate for each locus ranges from 0.44 (A10a) to 0.88 (A120a), and the across loci average is 0.74, suggesting that this set of loci is informative for population analyses.

Tests for Hardy-Weinberg equilibrium (HWE) using Fisher’s exact test with the sequential Bonferroni correction (Rice 1989) revealed that the populations conformed to Hardy–Weinberg equilibrium (HWE) at most loci. The very few observed locus/populations combinations that were not in HWE were not concentrated at any locus or in any population. Significant linkage disequilibrium was not detected between genotypes at the ten loci. As no evidence of linkage disequilibrium between loci was assessed, these 10 loci can be considered to be independent.

Population variability and differentiation

An estimate of variability distribution in and among the six tested populations (AMOVA) indicates that 90% of the variation occurs within populations while only about 10% of

| Locus  | na  | Min–Max | PIC  |
|--------|-----|---------|------|
| D4a    | 7   | 2–5     | 0.56 |
| D105a  | 15  | 5–11    | 0.72 |
| A7a    | 13  | 7–11    | 0.83 |
| A112a  | 18  | 8–12    | 0.82 |
| A115a  | 14  | 7–12    | 0.80 |
| A120a  | 15  | 7–13    | 0.88 |
| A122a  | 12  | 5–9     | 0.74 |
| A17a   | 11  | 6–9     | 0.77 |
| A10a   | 13  | 2–11    | 0.44 |
| C103a  | 11  | 7–9     | 0.80 |
| Mean   | 12.9| 5.6–10.2| 0.74 |

na, mean number of alleles; Min–Max, minimum and maximum number of alleles; PIC, polymorphic information content.
**Table 3.** Genetic variability of wild populations of *Anastrepha fraterculus* from different geographical regions in Brazil estimated using 10 SSRs.

| Population           | na   | He   | Ho   | $F_{IS}$ |
|----------------------|------|------|------|---------|
| Una-BA               | 7,7  | 0,63 | 0,54 | 0,14    |
| Porto Seguro-BA      | 8,2  | 0,70 | 0,67 | 0,02    |
| Monte Alegre-RN      | 7,5  | 0,68 | 0,57 | 0,20    |
| São Mateus-ES        | 6,3  | 0,66 | 0,66 | 0,08    |
| Campos do Jordão-SP  | 8,3  | 0,71 | 0,61 | 0,13    |
| Vacaria-RS           | 8,5  | 0,72 | 0,62 | 0,12    |

na, mean number of alleles; He, expected heterozygosity; Ho, observed heterozygosity; $F_{IS}$, fixation index.

**Table 4.** Spatial Analysis of Molecular Variance (SAMOVA) for different population partitions.

| Number of groups ($K$) | $F_{CT}$ | $P$       | Population partition                                      |
|------------------------|----------|-----------|----------------------------------------------------------|
| 2                      | 0.195    | 0.062     | (Una, Porto Seguro, Monte Alegre, São Mateus), (Campos do Jordão, Vacaria) |
| 3                      | 0.182    | 0.015     | (Una, Porto Seguro, São Mateus), (Monte Alegre), (Campos do Jordão, Vacaria) |
| 4                      | 0.190    | 0.050     | (Una, Porto Seguro, São Mateus), (Monte Alegre), (Campos do Jordão), (Vacaria) |
| 5                      | 0.196    | 0.068     | (Una, Porto Seguro), (São Mateus), (Monte Alegre), (Campos do Jordão), (Vacaria) |

Total variation is detected among populations. Indeed as shown in Table 3, the intrapopulation genetic variability is similar across the six samples. As a second step, the simulated annealing approach based on the SAMOVA algorithm was applied to identify the presence of genetically homogeneous groups across the considered Brazilian populations. For this, the spatial analyses of molecular variance was performed without constraints for geographic composition of the groups. As observed in Table 4 no great differences in the $F_{CT}$ values were observed when we increased the group number ($K$): four of the five simulated groupings (2, 4, and 5) produced non-significant $F_{CT}$ values. This implies that the molecular variance due to differences between populations within each group is weak, confirming the AMOVA data. Only in one case did we observe an $F_{CT}$ estimate which statistically maximized the differences between groups. This is the grouping configuration which splits the six populations into three groups: i) Una, Porto Seguro and São Mateus, ii) Monte Alegre, iii) Campos do Jordão and Vacaria, ($F_{CT} = 0.182$, $P = 0.015$). These data indicate that the mountain populations of Campos do Jordão (state of São Paulo) and Vacaria (state of Rio Grande do Sul) are genetically homogeneous and differentiated from the group of the three coastal populations (Una, Porto Seguro and São Mateus), and from the other more distant coastal population of Monte Alegre. The pairwise $F_{ST}$ estimates (Table 5) confirm the presence of differentiation between the group of the two mountain populations and the coastal populations which in turn share a certain degree of genetic relatedness. Principal Coordinate Analysis (PCoA) was performed to better clarify the genetic relations among the six populations. The first two axes explain a relatively high amount of the genetic variation (88%). The first axis (77.91%) separates Campos do Jordão and Vacaria from the other...
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The second axis (10.39%) mainly differentiates São Mateus from the group of Una, Porto Seguro, and Monte Alegre, but also Campos do Jordão from Vacaria. It is interesting that there is a certain correspondence of the genetic grouping with the known morphotype classification (Hernández-Ortiz et al. 2012). The molecular variance represented by the first axis separates the populations on the basis of both geographical distance and altitude. On this basis, we attempted to disentangle the effect of geographical distance and altitude on the genetic differentiation of Campos do Jordão and Vacaria. The plots in

**Table 5.** Pairwise-$F_{ST}$ values among 6 population samples of *Anastrepha fraterculus* as derived from Microsatellite Analyser (Dieringer and Schlotterer 2003).

|                  | Una-BA | Porto Seguro-BA | Monte Alegre-RN | São Mateus-ES | Campos do Jordão-SP | Vacaria-RS |
|------------------|--------|----------------|----------------|---------------|---------------------|------------|
| Una-BA           |        |                |                |               |                     |            |
| Porto Seguro-BA  | 0.015  |                |                |               |                     |            |
| Monte Alegre-RN  | 0.020  | 0.012  ns       |                |               |                     |            |
| São Mateus-ES    | 0.031  | 0.031           | 0.042          |               |                     |            |
| Campos do Jordão-SP | 0.163 | 0.127           | 0.130          | 0.160         |                     | 0.038      |
| Vacaria-RS       | 0.165  | 0.126           | 0.132          | 0.175         | 0.038               |            |

Some of the values are not significantly different from zero at $P > 0.05$ (ns)

**Figure 2.** Two-dimensional plot of Principal Coordinate Analysis (PCoA) based on similarity matrix derived from *Anastrepha fraterculus* microsatellites data. The Morphotype classification (Hernández-Ortiz et al. 2012), relative to each sample is also reported.
Figure 3. Correlation of $F_{ST}$ values with the geographic distances (upper plot) and altitude differences (bottom plot) among the 6 Brazilian samples of *Anastrepha fraterculus*. 
Figure 3 show the correlations between pairwise $F_{ST}$ values with the differences in altitude (Figure 3A) and the geographic distance (Figure 3B), respectively. The correlation results clearly indicate that the difference in altitude has a greater impact ($r^2 = 0.76$ vs $r^2 = 0.28$) on genetic differentiation of mountain Vacaria and Campos do Jordão, than on the remaining samples belonging to the coastal plain populations.

**Discussion**

With this paper we initiated, in the polymorphic *A. fraterculus* complex, an analysis of the underpinning genetic architecture and its interaction with correlated ecological, biological and morphological traits. In this context, we have used microsatellite markers to perform a genetic analysis of populations from a complex ecological area such as Brazil.

Although the chromosomal location of the considered SSR loci remains unknown, the assessed linkage equilibrium between them suggests they are statistically independent and that their variability patterns might reflect genome-wide patterns across populations. The six considered ecogeographic populations are here represented by highly polymorphic samples, which reflect a high degree of intrapopulation genetic diversity. Indeed only 10% of the total variability (AMOVA) is represented by the differences between the six geographic populations, while greater variability was found within populations.

The spatial analysis of genetic diversity indicates that the levels of diversity among the six populations vary significantly on an eco-geographical basis, as indicated by SAMOVA and PCoA data. More than by geographical distance, the genetic differentiation is influenced by altitude. The multivariate analysis of ten microsatellites depicts a structural pattern, which clearly separates populations on climatic distribution both on latitudinal and altitudinal basis. Particularly, altitude seems to represent a differentiating adaptation, as the main genetic differentiation is that detected between the populations present at higher altitudes (Campos de Jordão and Vacaria) and those populations from sea level. Genetic divergence between populations from low and high altitude areas has already been observed for populations within the *A. fraterculus* complex using isozymes (Morgante et al. 1980, Steck 1991, Selivon et al. 2005) and mtDNA (Steck and Sheppard 1993, Santos 1999, McPheron et al. 1999, Smith-Caldas et al. 2001, Barr et al. 2005). Steck (1991) concluded that the strong differences in allelic frequency between lowland and Andean populations of *A. fraterculus* in Venezuela was due to the fact that they actually represent two genetically distinct species albeit morphologically indistinguishable. These allopatric populations can be subject to divergent selection in response to ecological factors that change over large geographical scales such as altitude. Altitude may act as a barrier to gene flow as levels of life history divergence between high- and low-altitude populations can be correlated with levels of post-zygotic reproductive isolation (Orr and Smith 1998).
One interesting observation, which arises from our data, is that the observed structure of Brazilian populations is entangled with the presence of morphotypes. The actual number of these entities and their respective geographic range are questions that remain to be further elucidated. At the moment three different morphotypes are identified in Brazil (Hernández-Ortíz et al. 2012). As it appears clearly, the PCoA analysis depicts a genetic differentiative pattern that overlaps with the distribution of the known morphotypes. Now the open questions are: 1) is the observed population differentiation contributing to the underpinning genetic architecture of the morphotypes associated to these populations? and 2) do morphotypes track environmental variability? In retrospect, a further aim is to clarify the evolutionary relationships between populations, ecotypes, and morphotypes.

**Conclusion**

The population genetic approach, in addition to improving our knowledge of the underpinning genetic architecture of the *A. fraterculus* complex, is also important from an applied perspective. The overall level of genetic variability and the presence of differentiation that we detected among the Brazilian populations of *A. fraterculus* constitute an important contribution for any potential future application of SIT for the control of populations of this fruit fly pest in Brazil.

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