

Unrelated secondary reproductives in the neotropical termite *Silvestritermes euamignathus* (Isoptera: Termitidae)

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Abstract A termite colony is usually founded by a pair of alates, the primary reproductives, which produce all the nestmates. In some species, secondary reproductives appear to either replace the primaries or supplement colony reproduction. In termites, secondary reproductives are generally ergatoids derived from workers or nymphoids derived from nymphs. *Silvestritermes euamignathus* is a termite species that forms multiple nymphoid reproductives, and to date it was hypothesized that these secondary reproductives were the progeny of the primary founding reproductives. We developed markers for 12 microsatellite loci and used COI mitochondrial DNA (mtDNA) to genotype 59 nymphoid neotenics found in a colony of *S. euamignathus* to test this hypothesis. Our results showed that nymphoids of *S. euamignathus* are not all siblings. The microsatellite analysis suggests that the secondary reproductives derived from a minimum of four different pairs of reproductives belonging to at least two different matriline. This is the first record of non-sibling secondary reproductives

occupying the same nest in a higher termite. These unrelated reproductives might be the result of either pleometrotic colony foundation or colony fusion.

Keywords Microsatellites · mtDNA · COI · Genotyping · Neotenics

Introduction

Termites exhibit one of the most fascinating reproductive systems among the social insects. These insects present an incredible plasticity of castes participating in reproduction, from regular primary reproductives derived from alates to ergatoid reproductives derived from workers (Noirot 1956; Myles 1999), reproductive soldiers in *Zootermopsis* (Thorne et al. 2003), nymphoid neotenics derived from nymphs (Noirot 1956; Myles 1999), and the parthenogenetically-produced nymphoids in lower and higher termites, which succeed the queen (Matsuura et al. 2009; Vargo et al. 2012; Luchetti et al. 2013; Fougeyrollas et al. 2015). Primary reproductives are winged imagos and the ancestral reproductive forms, whereas ergatoids and nymphoids are, respectively, apterous and brachypterous neotenics, a novelty of termites.

According to the origin, relatedness, and number of active reproductives, termite colonies are classified as simple families, extended families, and mixed families (reviewed in Vargo and Husseneder 2011). Simple families are colonies headed by a regular monogamous pair, the royal couple, whereas extended and mixed families present multiple reproductives. When multiple neotenics appear to supplement or replace the founding queen, we classify such colonies as extended families, because both primary queen and neotenics contribute to the offspring, and all the individuals, with exception of the king, share the queen's mitochondrial DNA (mtDNA)

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haplotype and exhibit no more than four alleles at a single microsatellite locus. Families are extended even when the primary queen is already dead and all offspring is produced only by the neotenics.

When multiple female alates co-found a colony, we have a case of pleometrotic polygyny (Atkinson and Adams 1997; Hartke and Rosengaus 2013), and the colony is classified as a mixed family. Mixed families can also result from colony fusion (DeHeer and Vargo 2004, 2008). The offspring of a mixed family exhibit multiple allele combinations, and frequently more than four alleles at one or more microsatellite loci. Mixed families formed either by unrelated female alates or by colony fusion may also exhibit different matriline, as occurs in *Nasutitermes corniger* (Atkinson and Adams 1997), *Macrotermes michaelseni* (Hacker et al. 2005), and *Reticulitermes flavipes* (DeHeer and Vargo 2004, 2008). In both extended and mixed families, the multiple females may be accompanied by multiple males, because polyandry is often associated with polygyny in termites (Roisin and Pasteels 1985).

Silvestritermes euamignathus is a neotropical higher termite species that occurs predominantly in open vegetation, from southern Brazil to northern Venezuela (Rocha et al. 2012). This species presents nasute-mandibulate soldiers and builds epigeal nests of variable forms, most commonly exhibiting a dome form (Mathews 1977; Coles-de-Negret and Redford 1982). Replacement reproductives are relatively rare in higher termites when compared to lower termites (Miller 1969; Noirot 1969; Myles 1999), and genetic studies analyzing these individuals are equally scarce in higher termites. Costa-Leonardo et al. (1996) found multiple male and female nymphoid reproductives in one nest of this species without either primary reproductives and in two other nests containing only the king. Later, the authors were able to artificially reproduce this scenario by experimentally orphaning natural colonies (Costa-Leonardo et al. 1998). In their study, Costa-Leonardo et al. (1998) stated that most nests of *S. euamignathus* presented the royal couple and neotenics were absent. Morphological analysis of the neotenics suggested their active participation in reproduction (Costa-Leonardo et al. 1999), and it was hypothesized that the naturally occurring secondary reproductives found by Costa-Leonardo et al. (1996) were full siblings that differentiated upon the death of the primary queen. The occurrence of multiple neotenics in colonies of *S. euamignathus* makes this termite species a good candidate to investigate the genetic relationship within these individuals.

In this study, we investigated the relationship of multiple nymphoid neotenics found in a colony of *S. euamignathus*. We developed markers for 12 microsatellite loci to assess the genetic variation within the reproductives, and used COI mtDNA genotyping as a supplemental technique to determine the number of matriline present in a colony. We tested the

hypothesis by Costa-Leonardo et al. (1996) about the relationship within neotenics, and where genetic diversity was higher than expected for simple families, the number of matriline was used as support to elucidate the mechanism involved in the process.

Material and methods

Termites We collected five nests (colonies A–E) of *S. euamignathus* (Silvestri, 1901) in field areas at Rio Claro, SP, Brazil between September and November 2013. The nests were carried to the Laboratório de Cupins, UNESP campus, Rio Claro, SP. The nests were collected near to small fragments of natural Cerrado vegetation bordered by *Eucalyptus* plantations, frequently occurring in small ravines along the roads, or leaning tree trunks, but always easily detachable. They were collected along a range of 3 km distance (the highest distance between colonies A and E), with the smallest distance of 50 m apart between colonies B and C. Samples of all castes and all of the reproductives found were collected. The samples were stored in 95 % ethanol and sent to the Department of Entomology, North Carolina State University, Raleigh, NC, USA. One of these nests (colony D) contained 59 nymphoid neotenics: 27 males and 32 females, two of which being darker reproductives similar to alates in pigmentation, but with short wingbuds as in all the other neotenics (Fig. 1), whereas no reproductives were found in the other four colonies. All the nests shared the same shape and structure, the colonies were mature and exhibited nymphs or alates, but only the colony D containing the neotenics exhibited nymphs from all instars at the date of the collection.

Microsatellite marker development

DNA extraction for sequencing We extracted DNA from the heads of seven workers of *S. euamignathus* using a DNeasy

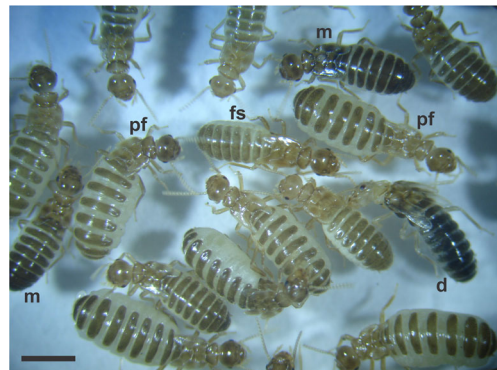


Fig. 1 Multiple nymphoid neotenics of *S. euamignathus* with differences in physogastry. *pf* physogastric females, *fs* females with a swollen abdomen, *m* male reproductives, *d* darker reproductives. Bar = 500 μ m

Blood & Tissue kit (Qiagen®) following the manufacturer's instructions. We used only termite heads to avoid contamination with the gut material and we pooled multiple individuals to obtain sufficient quantity of DNA for library preparation and sequencing.

DNA sequencing The Illumina MiSeq Reagent Kit V3 was used for the library preparation (300x300 Paired-end reads) and sequencing was performed using the Illumina MiSeq Sequencer. The sequencing generated approximately 22 M 300-bp paired-end reads, which were de novo assembled using the program SparseAssembler (Ye et al. 2012). The output product consisted of consensus regions of DNA (contigs), which were used to screen for the microsatellites.

Selection of microsatellite markers The contigs were analyzed using the Msatcommander® program for detecting microsatellite loci. The search parameter was set for trinucleotide microsatellite loci, with output files showing the repeats. We designed 54 primer pairs for the sequences containing microsatellite loci using Primer 3 v. 4.0.0 software (Koressaar and Remm 2007; Untergrasser et al. 2012). An M13 oligonucleotide fragment (CACGACGTTGTAACGAC) was added to the 5' end of each forward primer and was complementary to the LiCor IRDye-M13 fluorescent dye necessary for genotyping. The primers were optimized for PCR and the products of the reactions were analyzed on a Li-Cor 4300 automated DNA sequencer (Li-Cor, Inc., Lincoln, NE, USA). We used DNA from five samples, one individual from each colony, to select 12 polymorphic microsatellite markers for use in this study (Table 1). All the primers had an annealing temperature at 55 °C and we were not successful in combining different loci in multiplex reactions.

Microsatellite genotyping A total of 100 workers, 20 from each colony, and the 59 neotenic reproductives found in colony D were genotyped for the 12 selected loci. PCR reactions were carried out in 6- μ L volumes, each containing 7 ng of template DNA, 1 \times PCR buffer, 1.9 mM MgCl₂, 0.25 mM dNTPs, 1 pmol each primer (forward and reverse), 0.03 pmol M13F-29/IRD800 IRDye tag (Li-Cor, Inc), 0.05 U *Taq* DNA Polymerase (*Bioline TaQ Polymerase BIO-21086*) and ddH₂O. PCR products were separated by electrophoresis on 6 % polyacrylamide gels run at a constant power of 40 W at 50 °C for 2 h. The gel images were analyzed using GeneProfiler software (Scanalytics, Inc) to determine the allele sizes. All the alleles were an extra 19 bp in length due to the M13 tail.

Data analysis The number of alleles per colony was obtained for each locus using the Excel add-in Microsatellite toolkit. We also estimated *F*-statistics and relatedness within colonies

using FSTAT (Goudet 2002). For the FSTAT analysis we used the workers' genotypes.

Mitochondrial DNA

We used the DNA of the microsatellite-genotyped samples to run preliminary tests using three mitochondrial DNA genes: COI, COII and 16S. We used one sample from each colony to detect haplotype variation in the population. We selected the gene COI, which presented different haplotypes varying at eight positions. PCR reactions were carried out in 25- μ L volumes, each containing 50 ng of template DNA, 1 \times PCR buffer, 1.75 mM MgCl₂, 0.1 mM dNTPs, 0.1 pmol each primer (Forward—ATTCAACCAATCATAAAGATATNGG and Reverse—TATACTTCAGGGTGTCCGAAAAATCA), 0.07 U *Taq* DNA Polymerase (Bioline TaQ Polymerase BIO-21086) and ddH₂O. Cycling conditions were: initial denaturing at 95 °C for 3 min, followed by 35 cycles of 1 min denaturing at 95 °C, 1:30 min annealing at 45 °C and 2 min extension at 72 °C, and final step at 72 °C for 5 min. Based on the microsatellite analysis, we selected 40 individuals, five from each colony (A, B, C, and E) and 20 from colony D, composed of seven workers, six female nymphoids, five male nymphoids, and two darker reproductives to be sequenced. The selection was based on variation in the 12 microsatellite loci and the chosen samples presented different allelic combinations. The mitochondrial analysis was performed to supplement the microsatellite genotyping and to test whether the observed genetic diversity was result of multiple mating of a single queen.

Results

Microsatellite markers

Within the colonies genotyped, A, B, C, and E presented from one to four alleles per locus, which is in agreement with the occurrence of colonies initiated by single parents. Mean values of expected (HE) and observed (HO) heterozygosities were 0.6299 ± 0.0360 and 0.7808 ± 0.0255 , respectively. Colony D presented more than four alleles for nine out of the 12 loci (Table 2), indicating the presence of multiple unrelated reproductives in the colony. The neotenic presented a total of 72 different alleles among the 12 loci, and nine alleles were exclusively found among the females (Table 2). The different allelic combinations indicate the occurrence of at least three pairs of reproductives in the previous generation (with a maximum of four alleles per locus for each reproductive pair).

Overall, colonies were slightly inbred ($F_{IT} = 0.064 \pm 0.03$ (SE)), with moderate differentiation among them ($F_{CT} = 0.25 \pm 0.014$). Individuals were relatively outbred relative to others within their colonies ($F_{IC} = -0.248 \pm 0.028$),

Table 1 Characteristics of 12 microsatellite markers developed for *S. euamignathus*

Locus	Primer sequences	Repeat motif	N _A	Size (bp)	GenBank accession number
<i>Se1</i>	F: GGAGACAGTAACATGGTGCC R: AGAGGTAAGGCAGAATGGGA	(AAC) ₈	8	104–143	KU358999
<i>Se2</i>	F: ACTGGTCACTCTGCTTCACA R: ACCACGGATCTACATTCTTTCC	(AAT) ₈	12	224–272	KU359000
<i>Se3</i>	F: TGAGAAGCCATTGTCCACCA R: TGAGATCGAAATTGCCACAGT	(GTT) ₈	8	173–201	KU359001
<i>Se4</i>	F: AGTGCAGGGATGTCAAACCTG R: GGGCCTCATAACCTTGACCT	(ATC) ₉	7	163–258	KU359002
<i>Se5</i>	F: ACTGAACGAGTTGTCTGCAA R: GGTTTCTTCCATGACCACCA	(ATC) ₈	6	210–237	KU359003
<i>Se6</i>	F: GGAGGAGGACGAGGAAAAGG R: TGCCGTGTATGGAGTTCAGT	(AAG) ₁₁	12	199–307	KU359004
<i>Se7</i>	F: ACCCTGAAACCGAAAACGTT R: ACCTCTTCTTCAACAACAACA	(ACG) ₁₁	10	124–292	KU359005
<i>Se8</i>	F: TTGCTACTCCTGCCCTCATC R: GGGGATTACGCAAACCTTCG	(ATT) ₁₃	8	111–144	KU359006
<i>Se9</i>	F: GCTGAGAGAGTGGACTGGAC R: TTTCTCGGCTGCACTATTGT	(ATT) ₈	9	173–215	KU359007
<i>Se10</i>	F: CGGTGTGCATATTGTTGAGC R: TCTCCAGTGTGCTTGATACC	(ATC) ₉	4	129–138	KU359008
<i>Se11</i>	F: TGAACATGAACTCTGCAACTCA R: GGCCACTATCCCATTAAGCA	(AGC) ₈	6	197–215	KU359009
<i>Se12</i>	F: ACCCTGAAGCAAAGAAACCTC R: TCCACTCGACACCCTAGTTC	(ATT) ₈	13	144–213	KU359010

N_A is the number of alleles found in 159 individuals from 5 colonies for each locus

indicating few reproductives present in colonies on average. Finally, individuals within colonies were nearly as related as full siblings ($r=0.469\pm 0.019$). Together, these analyses indicate that most of the study colonies had a breeding structure close to a simple family, although none of the colonies was a

Table 2 Alleles per locus found in reproductives of colony D

Locus	N _A	Alleles
<i>Se1</i>	6	107*, 113, 125, 128, 131, and 134
<i>Se2</i>	10	224, 236, 239, 242, 245, 248, 254*, 260, 263, and 266
<i>Se3</i>	5	173, 176, 179, 185, and 194
<i>Se4</i>	5	163, 166, 175, 193, and 246
<i>Se5</i>	7	212, 215, 224*, 227*, 230, 233, and 236
<i>Se6</i>	8	199, 202, 205, 208, 220*, 223, 232, and 289
<i>Se7</i>	6	124, 136, 193, 202, 211, and 217
<i>Se8</i>	3	120, 123, and 141
<i>Se9</i>	6	173, 176, 179, 182, 200, and 206
<i>Se10</i>	4	129, 132, 135, and 138
<i>Se11</i>	3	200, 203, and 206*
<i>Se12</i>	9	156*, 162, 165*, 168, 171, 174, 180, 186, and 204*

N_A number of alleles

*alleles exclusively found among the female neotenic

simple family due to a small number of genotypes inconsistent with a single pair of parents. Colony D with 59 neotenic and an F_{IC} value of -0.0195 was an obvious exception to the pattern seen in the other four colonies.

The genotyped neotenic differed in size, with some females presenting advanced physogastry while others showed only a slightly swollen abdomen (Fig. 1). The allelic combination of these individuals did not differ from the physogastric neotenic, and this difference in size might be related to the nymphal instar of their origin.

Mitochondrial DNA

The fragment of COI region amplified was 620 bp long, and the variation in the haplotypes occurred at positions 29, 71, 140, 248, 464, 518, 587, and 590 in the gene sequences. We found three different haplotypes for COI within the five colonies of *S. euamignathus*. Colonies A and C shared haplotype I, whereas colonies B, D, and E shared haplotype II. Colony D, which had the multiple reproductives, presented two different haplotypes, one shared with colonies B and E (haplotype II) and another unique haplotype (haplotype III). Haplotype variation is summarized in Table 3.

Table 3 Haplotype variation for COI found in the studied population of *S. euamignathus*

Colony	Base Pair Position								GeneBank Accession Number
	29	71	140	248	464	518	587	590	
A	T	T	C	T	C	T	G	T	KU521793
B	T	C	T	C	T	C	G	C	KU521794
C	T	T	C	T	C	T	G	T	KU521795
D	T	C	T	C	T	C	G	C	KU521796
	C	T	C	T	C	T	A	T	KU521791
E	T	C	T	C	T	C	G	C	KU521792

Haplotype I: light gray; haplotype II: white; haplotype III: dark gray

The occurrence of two haplotypes in colony D is indicative of at least two different females producing the neotenic. In addition, the occurrence of more than four alleles at specific loci, such as locus *Se2* where we found ten alleles within the neotenic, suggests at least three different couples sharing haplotype II in the parental generation (Fig. 2). Depending on the loci, some alleles were exclusively present within the individuals of one or the other haplotype (Table 4).

Discussion

Independent colony foundation by unrelated queens was reported for *N. corniger* by Atkinson and Adams (1997). In this species, secondary reproductives may arise from pleometrotic associations, and this system leads to polycalism, in which a single colony headed by multiple unrelated reproductives is widespread among different nests (“calies”). Colonies with related reproductives occurred in single nests, while unrelated reproductives within a colony are allocated in different “calies”. This later situation was described by Thorne (1982) as “voluntary” budding. In our case, a single nest of *S. euamignathus*, colony D, contained 59 secondary reproductives belonging to at least two different matriline. The results from the mtDNA analysis are in agreement with the microsatellite genotyping,

which indicated that the neotenic originated from four or more primary couples. Both subgroups of neotenic from colony D sharing one or the other mtDNA haplotype presented more than four microsatellite alleles at a single locus, which suggests the presence of non-sibling reproductives in the previous generation. Of the five colonies sampled in this study, colony D was alone in this regard. Admittedly, it is possible that colony D was atypical, resulting from a set of highly unique circumstances and does not represent a common occurrence in *S. euamignathus*. However, given that we sampled only a small number of colonies, this possibility seems remote and it is likely that colonies with unrelated neotenic occur with some frequency in nature, at least in some populations.

The current scenario allows us to raise three hypotheses. First, two different female alates each mated with more than one male. This hypothesis unites the occurrence of pleometrotic colony foundation with a polyandry mating system by the co-founding females. Although this hypothesis seems to be the less probable because of the absence of known polyandry in primary reproductives in termites (Hartke and Baer 2011), a recent study by Wu et al. (2013) found evidence of polyandry in neotenic in the termite *Reticulitermes labralis*. These authors showed that one female neotenic mated with at least two male neotenic, as the alleles present in the offspring did not correspond to either asexual reproduction or

Fig. 2 Projection of the parental generation considering the two mtDNA haplotypes and the allelic variation (A^{1-10}) found within the progeny at a single locus (*Se2*)

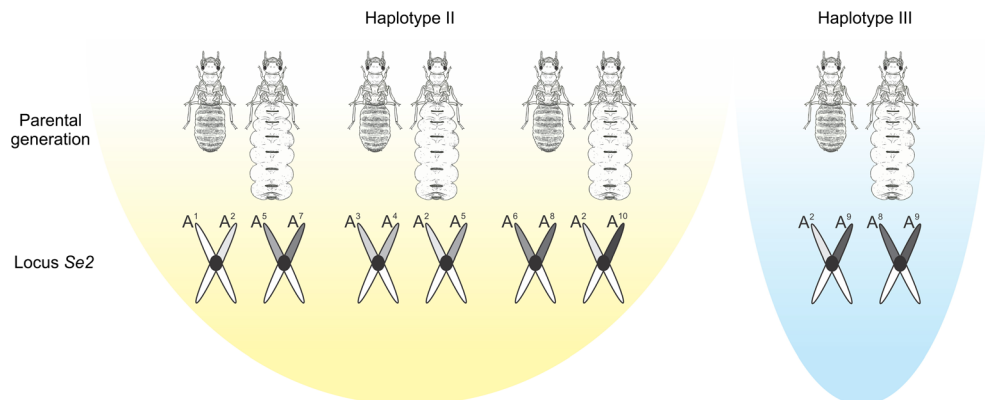


Table 4 Microsatellite loci genotypes of the 13 neoteniics used for mtDNA haplotype analysis

Neoteniics	Microsatellite loci genotypes																							
	Se1		Se2		Se3		Se4		Se5		Se6		Se7		Se8		Se9		Se10		Se11		Se12	
	A	A	A	A	A	A	A	A	A	A	A	A	A	A	A	A	A	A	A	A	A	A	A	A
	1	2	1	2	1	2	1	2	1	2	1	2	1	2	1	2	1	2	1	2	1	2	1	2
FN 01	1	1	2	2	1	1	1	1	2	2	2	1	2	1	1	1	2	2	1	1	2	2	1	1
	3	3	6	3	7	7	9	6	3	1	2	9	1	9	2	2	0	0	3	2	0	0	8	7
	1	1	0	6	9	6	3	3	3	5	3	9	7	3	3	3	0	0	5	9	0	0	6	4
FN 10	1	1	2	2	1	1	1	1	2	2	2	2	1	1	1	1	2	1	1	1	2	2	1	1
	3	0	6	4	7	7	6	6	3	2	3	3	2	2	2	2	0	8	3	3	0	0	8	7
	4	7	6	8	9	6	6	3	6	4	2	2	4	4	0	0	0	2	5	2	0	0	0	1
FN 11	1	1	2	2	1	1	2	1	2	2	2	2	2	2	1	1	2	1	1	1	2	2	1	1
	2	2	6	6	8	7	4	6	3	3	2	0	1	1	4	2	0	7	3	2	0	0	6	6
	8	8	3	3	5	3	6	3	6	0	3	8	7	7	1	0	6	9	8	9	6	0	8	2
FN 15	1	1	2	2	1	1	1	1	2	2	2	2	2	1	1	1	1	1	1	1	2	2	2	1
	3	0	5	4	7	7	7	6	2	1	8	3	1	2	2	2	8	7	3	3	0	0	0	5
	4	7	4	8	6	6	5	3	7	2	9	2	7	4	0	0	2	6	5	2	3	0	4	6
FN 22	1	1	2	2	1	1	1	1	2	2	2	1	2	2	1	1	1	1	1	1	2	2	1	1
	3	2	6	3	9	7	9	6	3	3	0	9	1	0	2	2	7	7	2	2	0	0	7	7
	1	5	0	6	4	3	3	3	3	3	8	9	7	2	3	0	9	9	9	9	3	0	1	1
FN 26	1	1	2	2	1	1	1	1	2	2	2	2	2	1	1	1	1	1	1	1	2	2	1	1
	3	2	3	3	7	7	9	9	3	1	2	0	1	2	2	2	7	7	2	2	0	0	7	7
	1	5	6	6	9	9	3	3	3	5	0	8	7	4	0	0	9	9	9	9	0	0	4	1
DFN 31	1	1	2	2	1	1	1	1	2	2	2	1	2	1	1	1	1	1	1	1	2	2	1	1
	3	3	3	3	9	9	9	6	3	1	2	9	1	9	2	2	7	7	2	2	0	0	7	7
	1	1	6	6	4	4	3	3	0	5	3	9	7	3	3	3	9	9	9	9	3	0	1	1
DFN 32	1	1	2	2	1	1	1	1	2	2	2	2	1	1	1	1	1	1	1	1	2	2	1	1
	2	2	6	3	8	7	7	6	3	3	2	0	3	3	2	2	7	7	3	2	0	0	8	6
	8	5	0	6	5	9	5	3	0	0	3	2	6	6	0	0	9	9	8	9	3	0	6	2
MN 08	1	1	2	2	1	1	1	1	2	2	2	1	2	2	1	1	2	2	1	1	2	2	1	1
	3	2	4	2	7	7	9	6	1	1	0	9	0	0	2	2	0	0	2	2	0	0	7	7
	1	5	5	4	9	6	3	3	5	5	8	9	2	2	3	3	0	0	9	9	3	0	4	1
MN 12	1	1	2	2	1	1	1	1	2	2	2	2	1	1	1	1	2	1	1	1	2	2	1	1
	3	2	4	3	7	7	9	6	3	3	0	0	2	2	2	2	0	7	3	2	0	0	7	6
	1	5	2	6	6	3	3	3	3	0	8	2	4	4	3	0	0	9	5	9	3	0	4	8
MN 18	1	1	2	2	1	1	1	1	2	2	2	1	1	1	1	1	2	1	1	1	2	2	1	1
	3	2	6	3	7	7	9	7	3	1	2	9	9	3	2	2	0	7	2	2	0	0	8	7
	1	5	0	6	9	9	3	5	3	5	3	9	3	6	3	0	0	9	9	9	3	0	6	1
MN 19	1	1	2	2	1	1	1	1	2	2	2	2	2	2	1	1	2	1	1	1	2	2	1	1
	2	1	4	3	7	7	7	6	3	3	0	0	0	0	2	2	0	7	3	3	0	0	7	6
	5	3	5	9	9	9	5	3	3	3	5	2	2	2	3	0	0	3	8	5	3	0	4	2
MN 26	1	1	2	2	1	1	1	1	2	2	2	1	2	1	1	1	1	1	1	1	2	2	1	1
	3	2	3	2	7	7	9	6	1	1	0	9	0	9	2	2	7	7	2	2	0	0	8	7
	1	5	6	4	9	9	3	3	5	5	8	9	2	3	3	0	9	9	9	9	3	3	6	1

The two haplotypes found within the neoteniics from colony D is marked in *white* and *dark gray* in the neotenic code column. Alleles marked in *yellow* were exclusively found in individuals with mtDNA haplotype II (*white*) and those marked in *blue* in individuals with mtDNA haplotype III (*dark gray*). Alleles in bold were exclusively found in these individuals

A1 and A2 are alleles, FN female neoteniics, DFN darker female neoteniics, MN male neoteniics

sexual reproduction with a single male (simple family). As we did not have access to the parental generation because they were not present in the nest at the time of the collection, we cannot exclude this hypothesis.

The second hypothesis is that there is pleometrosis during colony foundation with multiple reproductive pairs, but only one male for each female. Pleometrotic associations were

already recorded for the higher termites *N. corniger* (Thorne 1982, 1984; Atkinson and Adams 1997) and *M. michaelsoni* (Hacker et al. 2005). In the latter species, the number of unrelated primary queens tends to decrease during the development of the colony, and few queens are likely to coexist in mature colonies. In this study, the genotyping results showed that at least four independent couples produced the 59

secondary reproductives found in colony D. Three or more unrelated couples belonging to one maternal lineage generated some of the neotenic reproductives, and one or more unrelated couples belonging to another maternal lineage generated the other neotenic.

The third hypothesis is based on mixed families arising through colony fusion. Colony fusion often leads to the death of primary reproductives and the inheritance of the colony by replacement reproductives. Some termite species, mostly lower termites, exhibit colony fusion (DeHeer and Vargo 2004, 2008; Guaraldo and Costa-Leonardo 2009; Luchetti et al. 2013). Although there is no record of colony fusion for *S. euamignathus*, this termite often occupies abandoned nests of other termite species and restructures them (Rocha et al. 2012). This behavior indicates that either abandoned nests are colonized by alates after swarming or colonies have a certain mobility. Whether colonies are mobile and occupy nests built by conspecifics is difficult to know, because an inquiline colony would not need to modify the structure of a nest constructed by conspecifics. Indeed, no differences in nest shape and structure were observed among the colonies used in this study. The unique difference observed within the colonies was in respect to the nymphs, because all instars of the post-embryonic development were found in the colony with neotenic. Matsuura and Nishida (2001) reported that colony fusion in the termite *Reticulitermes speratus* is influenced by nymph ratios between the colonies. Whether the plenty of nymphs in the colony D was the key factor for colony fusion or a result of that, this scenario represents a disequilibrium of the colony reproductive system and, as suggested by Costa-Leonardo et al. (1999), the occurrence of multiple secondary reproductives would be a transitional situation after the disappearance of primaries in *S. euamignathus*, which is probable in the case of colony fusion.

Multiple secondary reproductives were also found by Barbosa et al. (2012) in a nest of *Silvestritermes holmgreni* without the primaries, suggesting that the occurrence of replacement reproductives may be widespread in the genus *Silvestritermes*. The female nymphoids of *S. holmgreni* also varied in size, and these differences were based on the instar of the nymph that differentiated into neotenic (Barbosa et al. 2012). Different degrees of physogastry were also found within female neotenic in *S. euamignathus*, which seem to be associated with the nymphal instar that differentiated into neotenic (Costa-Leonardo et al. 1999, Haifig et al., unpublished). In addition, we also collected the two darker reproductives that resemble nymph-imago-neotenic intercastes or pseudoimagos differentiating at a very late stage of the imaginal development, which may have appeared in response to orphaning of the colony, such as observed by Roisin and Pasteels (1986a, b) for *Nasutitermes*. Based on morphological

characters, Rocha et al. (2012) pointed out in their taxonomic revision the close relationship between *S. euamignathus* and *S. holmgreni*, and the authors did not exclude the possibility of synonymy. As evidenced by morphology, the current scenario is still dubious and future genetic studies using the microsatellite markers developed here in colonies and populations of the genus *Silvestritermes* will provide a better understanding of the relationship among the species and the origin and relatedness of secondary reproductives in the group.

In sum, our results showed that colonies are often simple families in *S. euamignathus* and new colonies are likely to be founded by single couples. The occurrence of multiple reproductives in one of the colonies in this study, together with previous records of co-occurring multiple reproductives (Araujo 1958; Noirot 1956; Costa-Leonardo et al. 1996, 1998, 1999), indicates plasticity in the reproductive strategies in this termite species. The genetic analysis showed that the multiple secondary reproductives found within a colony were unrelated, indicating that they inherit a colony previously headed by unrelated reproductives. This is the first record of non-sibling neotenic reproductives in higher termites living within a single nest. Whether this reproductive condition is the result of pleometrotic colony foundation or colony fusion in *S. euamignathus* requires further investigation.

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