



# The mitochondrial genome of the stingless bee *Melipona bicolor* (Hymenoptera, Apidae, Meliponini): Sequence, gene organization and a unique tRNA translocation event conserved across the tribe Meliponini

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## Abstract

At present a complete mtDNA sequence has been reported for only two hymenopterans, the Old World honey bee, *Apis mellifera* and the sawfly *Perga condei*. Among the bee group, the tribe Meliponini (stingless bees) has some distinction due to its Pantropical distribution, great number of species and large importance as main pollinators in several ecosystems, including the Brazilian rain forest. However few molecular studies have been conducted on this group of bees and few sequence data from mitochondrial genomes have been described. In this project, we PCR amplified and sequenced 78% of the mitochondrial genome of the stingless bee *Melipona bicolor* (Apidae, Meliponini). The sequenced region contains all of the 13 mitochondrial protein-coding genes, 18 of 22 tRNA genes, and both rRNA genes (one of them was partially sequenced). We also report the genome organization (gene content and order), gene translation, genetic code, and other molecular features, such as base frequencies, codon usage, gene initiation and termination. We compare these characteristics of *M. bicolor* to those of the mitochondrial genome of *A. mellifera* and other insects. A highly biased A+T content is a typical characteristic of the *A. mellifera* mitochondrial genome and it was even more extreme in that of *M. bicolor*. Length and compositional differences between *M. bicolor* and *A. mellifera* genes were detected and the gene order was compared. Eleven tRNA gene translocations were observed between these two species. This latter finding was surprising, considering the taxonomic proximity of these two bee tribes. The tRNA<sup>Lys</sup> gene translocation was investigated within Meliponini and showed high conservation across the Pantropical range of the tribe.

*Key words:* *Melipona bicolor*, Meliponini, social insects, mitochondrial genome, gene rearrangements.

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## Introduction

In recent years a number of mitochondrial genomes have been completely sequenced, contributing to the knowledge of molecular features related to function and evolution of this peculiar genome (Boore, 1999). Its gene content is typically rather conserved in Metazoa (Boore *et al.*, 1995), with notable exceptions described in nematodes (Okimoto *et al.*, 1992), molluscs (Hoffmann *et al.*, 1992) and cnidarians (Beagley *et al.*, 1998). In general, mitochondrial DNA (mtDNA) contains genes for two ribosomal subunits (12S and 16S), 22 tRNA, and 13 proteins (three subunits of cytochrome c oxidase, cytochrome B, subunits

6 and 8 of ATP F0 synthase, and seven subunits of NADH dehydrogenase). There is also a non-coding A+T rich region that contains signaling elements for regulation of replication and transcription (Wolstenholme, 1992). Animal mitochondrial genome size is usually about 16 kb long, with few exceptions (Brown, 1985). Gene rearrangements within the mtDNA, formerly considered rare, have been described for a number of taxa, including bees (Mindell *et al.*, 1998; Dowton and Campbell, 2001; Shao *et al.*, 2003; Silvestre *et al.*, 2002; Silvestre and Arias, 2006).

One hundred sixteen complete arthropod mtDNA sequences, including 61 insect species, have been deposited in GenBank. In the described insect mtDNA genomes, A+T content is very high. Currently, complete mitochondrial genomes have been sequenced for just two hymenopteran species: the sawfly *Perga condei* (Castro and Dowton, 2005) and the honeybee *Apis mellifera* (Crozier and Cro-

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zier, 1993). The honeybee mtDNA presents the highest A+T bias (84.9%) known for insects. According to Crozier and Crozier (1993) the A+T bias is probably maintained due to a greater number of transversions over transitions.

Among bees, the tribe Meliponini, known as stingless bees, has gained some attention. The tribe has a wide geographic distribution, inhabits all tropical areas of the World (Michener, 2000), and includes the main pollinators in several neotropical ecosystems (Kerr *et al.*, 1996). In addition to their ecological importance, some species produce honey, pollen, wax and propolis that are commercially exploited (Nogueira-Neto, 1997). The species *Melipona bicolor* presents a very unique behavioral characteristic, polygyny, where several laying queens may cohabitate the nest for considerable time (Velthuis *et al.*, 2006). This species is distributed in the southern and southeastern Brazil, in the Atlantic rain forest. Nonetheless as this ecosystem has been severely reduced in size by human activity and only 7% of the original area remains, *M. bicolor* and other bees are considered endangered.

In previous studies we have characterized the mitochondrial genomes of several meliponine species through RFLP analysis (Francisco *et al.*, 2001; Weinlich *et al.*, 2004; Brito and Arias, 2005). Evidences of size difference, in comparison to those expected for *Apis mellifera*, were obtained for some mitochondrial regions after PCR amplification. Those data were interpreted as indirect evidence of gene order or content changes. Later these size differences, relative to *A. mellifera*, were confirmed by sequencing (Silvestre *et al.*, 2002; Silvestre and Arias, 2006), and all involved tRNA gene rearrangements. Eleven tRNA translocations were mapped between *A. mellifera* and *M. bicolor* (Arias *et al.*, 2006), so far the highest translocation number verified between species belonging to the same taxonomic subfamily.

Mitochondrial gene order has been considered a molecular class very promising for phylogenetic studies, especially among major taxonomic groups (Boore, 1999). tRNA rearrangements are highly reliable as evolutionary markers, neutral and less prone to homoplasy (Boore and Brown, 1998). We previously sequenced a portion of the mitochondrial genome of *M. bicolor*, and reported that there were 11 tRNA gene rearrangements when this genome was compared with that of *Apis mellifera* (Silvestre *et al.*, 2002; Silvestre and Arias, 2006). However, in that report we did not extensively characterize the *M. bicolor* mitochondrial genome, focusing our observations on genome organization. In the present study, we present a thorough characterization of the *M. bicolor* mitochondrial genome, with respect to codon usage, base frequencies, gene initiation and termination, and tRNA structure. In addition, we examine the evolutionary origin of one of the tRNA gene rearrangements identified in the previous study, the translocation of the tRNA<sup>Lys</sup> gene. We survey a broad range of Meliponini to more accurately determine the evolutionary origin of this translocation.

## Material and Methods

### mtDNA sequence analysis

Individuals of *M. bicolor* were collected from a single monogynic colony maintained at the Laboratório de Abelhas, Departamento de Ecologia, IB-USP, São Paulo. Total genomic DNA was extracted using the TNE-protocol (Sheppard and McPheron, 1991). The experimental approach to obtain the whole mtDNA molecule consisted of the amplification of small and overlapped fragments. Thus aliquots of TNE extraction were used as template for PCR reactions, with *Taq* DNA polymerase (Invitrogen). We initially used the following conditions: denaturation at 94 °C/5 min, followed by 35 cycles of 94 °C/60 s, 42 °C/80 s and 64 °C/120 s. An additional final extension step of 64 °C for 10 min was performed. The annealing temperature was optimized when necessary. The primers were derived from *A. mellifera* (Hall and Smith, 1991; Arias *et al.*, 2008), *M. bicolor* (designed in our laboratory), and other organisms (Simon *et al.*, 1994) (Table 1).

The PCR products were cloned into pGEM T-Easy Vector (Promega). At least two clones of each region were sequenced on both directions using Thermo Sequenase Dye Terminator (Amersham Life Science) or Big Dye Terminator (Applied Biosystems). Samples were analyzed on ABI-PRISM 310 and 3100 automated sequencers (Applied Biosystems).

Electropherograms were checked by eye on TraceViewer 2.0.1, and the sequences were assembled manually through GeneRunner 3.00 program (Hastings Software). Automated alignments were obtained using DAMBE (Xia, 2000), employing the CLUSTALW algorithm. Codon usage was analyzed with the software CODONTREE. BLAST searches at National Center for Biotechnology Information (NCBI) were used to verify the similarity between our sequences and those from other insect mitochondrial genomes. Transfer RNA genes were identified and the secondary structures were inferred by the software tRNA-Scan (Lowe and Eddy, 1997).

A single sequence of 14,422 bp was assembled and deposited at GenBank database under the accession number AF466146, which has also a Genome accession number, NC\_004529.

### KD tRNA cluster analysis

Fourteen Meliponini species (Table 2) were selected for the KD tRNA cluster sequencing. The species analyzed, collected in four continents, may be considered representative of the Pantropical geographic distribution of the tribe. The KD region was amplified using the pair of primers Cox 2/ Atp8 and the PCR conditions described in Castro *et al.* (2002). The computational analyses were as described above.

**Table 1** - Set of primers utilized on PCR amplification of *M. bicolor* mtDNA: positioning (relative to mitochondrial genes), strand, sequences and references.

Primer	Position (gene)	Strand	Sequence (5' - 3')	Reference
*11781F	CytB	<	TTTAAAATTATCTGGATCTC	(this work)
*16SF	16S	<	CACCTGTTTATCAAAAACATGTCC	Hall and Smith, 1991
*16SR	16S	>	CGTCGATTTGAACCAAAATCATG	Hall and Smith, 1991
*8321R	ND5	>	TTATATATCTAATTCTAT	(this work)
*8467F	ND5	<	GGAATTTTTTTTTGAATGAAA	(this work)
*C1-J-1718	COI	>	GGATCACCTGATATAGCATTCCC	Simon <i>et al.</i> , 1994
*C2-J-3696	COII	>	GAAATTTGGAGCAAATCATAG	Simon <i>et al.</i> , 1994
*C2-N-3661	COII	<	CCACAAAATTTCTGAACATTGACCA	Simon <i>et al.</i> , 1994
*C3-N-5460	COIII	<	TCAACAAAAGTGTCAAGTATCA	Simon <i>et al.</i> , 1994
*CB-J-10933	CytB	>	TATGTACTACCATGAGGACAAAATATC	Simon <i>et al.</i> , 1994
*N1-J-12585	ND1	>	GGTCCCTACGAATTTGAATATATCCT	Simon <i>et al.</i> , 1994
*N1-N-12595	ND1	<	GTAGCATTTTTAACTTTATTAGAACG	Simon <i>et al.</i> , 1994
*ND4F	ND4L	<	ATAAATTATGAACCTGGTCATCA	(this work)
*Mbi 18	ND3	<	GAACTATCAATTTGATATTG	(this work)
*Mbi 29	ND4	>	CGAAATTCACCATAACCTC	(this work)
*SR-N-14588	12S	<	AAACTAGGATTAGATACCCTATTAT	Simon <i>et al.</i> , 1994
*TM-J-206	RNA <sup>Met</sup>	>	GCTAAATAGCTAACAGGTTTCAT	Simon <i>et al.</i> , 1994
5612R	COIII	>	GAAATTAATATAACATGACCACC	Arias unpublished data
AMB 1	RNA <sup>Ile</sup>	>	TGATAAAAAGAAATATTTTGA	Arias unpublished data
AMB 3	RNA <sup>Trp</sup>	>	TTTAAAAACTATTAATCTTC	Arias unpublished data
AMB 4	COI	<	GAAAGTTAGATTTACTCC	Arias unpublished data
L2Eco	ND2	<	TTTTAGAATTCTGAAAATAATATAAAATTG	(this work)
Mel 2	ATPase 8	<	TGGAAAAATAATAATTG	(this work)
MtD 35Eco	12S	>	ATTCGAATTCAACTGAACGGGCAATTTGT	(this work)
Mbi 02	COI	>	GATCCAATAGGAGGAGGAG	(this work)
Mbi 03	COII	<	GGTATTAACGTTCAAATAATTTTC	(this work)
Mbi 04	CytB	>	CAATTCCAAATAAAATTAGGAGG	(this work)
Mbi 05a	ND1	<	TTAATCGAGTTCCATTG	(this work)
Mbi 06	16S	>	CAATTGATTATGCTACCTTTG	(this work)
Mbi 09	COI	>	GATTTCCATTAATTTTCAGG	(this work)
Mbi 10	ATPase 8	>	TTGACTTATTCCTTCATG	(this work)
Mbi 11	COIII	<	CGGATAATATCTCGAAATC	(this work)
Mbi 12	ND1	>	CTAATTCTGATTCTCCCTC	(this work)
Mbi 13	16S	>	CCGTGATACAAAAGGTAC	(this work)
Mbi 14	COII	>	TAATATGGCAGATTAAGTGC	(this work)
Mbi 16	COIII	>	TAAAAAATAGTCCCGACC	(this work)
Mbi 19	ND3	<	TGGGATTGAATCCATATTC	(this work)
Mbi 21	COII	>	CTATTA AAAACAATTGGTCATC	(this work)
Mbi 22	COI	<	GATGTAAAAATATGCTCGTG	(this work)
Mbi 23	ND5	<	TCGATTCGGAGATAGAGG	(this work)
Mbi 24	COIII	>	GATTTGAGATATTATCCG	(this work)
Mbi 25	RNA <sup>Arg</sup>	>	GATTTATGTTTAAAGTCGA	(this work)
Mbi 27	16S	<	GTTTAAAGTAATGTAAATG	(this work)
Mbi 30	ND1	<	TCGAGTTCCATTGATTT	(this work)
Mbi 31	ND5	>	TCGAGTTCCATTGATTT	(this work)
Mbi 32	CytB	<	AATGCAGTTGCTATTGATA	(this work)
Mbi 33	ND4L	>	TTTTGATGGACCCAAATTC	(this work)
Mbi 34	ND5	<	TCTACATTAAGACAATTAGG	(this work)
Mbi 35	ND5	>	CATTATAAAAACCTAATTGTC	(this work)
Mbi 36	ND4	>	TTCCATGTTCAACAAATCT	(this work)
Mbi 37	ND4	<	TATATTTAATTCATGGTTG	(this work)
Mbi 38	RNA <sup>Pro</sup>	>	TTAAATTATCTACAATCCC	(this work)
Mbi 39	CytB	<	TATTGATAGGTGTAGGAAG	(this work)
Mbi 40	ND2	>	CTATATCATCTATTTTCATC	(this work)
Mbi 41	COI	<	CATATAAAAGATTA AAAATTC	(this work)
Mbi 43	ND2	>	TAAAACAATTTATCATATTTTG	(this work)
Mbi 44	ND2	<	TAAATAATAATTTTATATG	(this work)
Mbi 47	ND6	<	TGAAAATAAAAATAATTTGG	(this work)
TPheF	RNA <sup>Phe</sup>	<	GCGTAATATTGAAAATATTAATGA	(this work)

\*Primers initially used to amplify longer amplicons, covering most of the genome. These regions were re-amplified in smaller fragments to continue the sequencing process.

**Table 2** - Meliponini species and countries of origin.

Species	Country
<i>Austroplebeia australis</i>	Australia
<i>Austroplebeia symei</i>	Australia
<i>Trigona carbonaria</i>	Australia
<i>Trigona hockingsi</i>	Australia
<i>Lestrimellita limao</i>	Brazil
<i>Plebeia remota</i>	Brazil
<i>Scaptotrigona xanthotricha</i>	Brazil
<i>Schwarziana quadripunctata</i>	Brazil
<i>Tetragonisca angustula</i>	Brazil
<i>Dactylurina</i> sp	Ghana
<i>Liotrigona</i> sp	Ghana
<i>Heterotrigona iridipennis</i>	India
<i>Trigona doipaensis</i>	Thailand
<i>Trigona flavivasis</i>	Thailand

## Results and Discussion

### General features of the genome and gene content

The size of the *M. bicolor* mtDNA had been previously estimated to be 18,500 bp by RFLP analyses (Weinlich *et al.*, 2004). However, this total size could not be fully confirmed by sequencing, we were unable to clone the 4,100 bp fragment containing the control region and its adjacent region. We analyzed a continuous fragment of 14,422 bp, or about 78% of the estimated size. This fragment contains the 13 protein-coding genes, 18 of 22 tRNA genes and the two rRNA genes (complete sequence was obtained for the large subunit - 16S, and partial sequence for the small subunit - 12S) (Figure 1). We detected five overlapping regions between genes, three of them including genes that were on the same strand (Table 3). In total, 30 bp were involved.

Seventeen non-coding regions were detected, with sizes ranging from one to 173 bp, totaling 486 bp (Table 4). Considering the same portion of *A. mellifera* mtDNA, excluding the hypervariable COI-COII intergenic region and the control region, the number of non-coding nucleotides is

**Table 3** - Overlapping regions between mitochondrial genes of *M. bicolor*: involved genes, overlap size (bp) and coding strand.

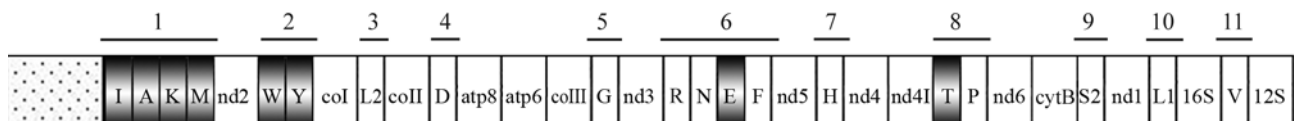
Genes	Size (bp)	Strand
ATP8/ATP6	10	+/+
tRNA <sup>Glu</sup> /tRNA <sup>Phe</sup>	6	+/-
tRNA <sup>His</sup> /ND4	6	-/-
ND1/tRNA <sup>Leu(L1)</sup>	6	-/-
tRNA <sup>Ser(S2)</sup> /ND1	2	+/-
Total	30	

**Table 4** - Non-coding regions between mitochondrial genes of *M. bicolor*: flanking genes and size (bp).

Genes	Size (bp)
tRNA <sup>Met</sup> /ND2	173
ND6/CytB	94
tRNA <sup>Asn</sup> /tRNA <sup>Glu</sup>	58
tRNA <sup>Tyr</sup> /COI	58
tRNA <sup>Pro</sup> /ND6	23
ATP6/COIII	15
COII/tRNA <sup>Asp</sup>	12
ND4/ND4L	10
COIII/tRNA <sup>Gly</sup>	9
CytB/tRNA <sup>Ser(S2)</sup>	9
tRNA <sup>Lys</sup> /tRNA <sup>Met</sup>	6
ND2/tRNA <sup>Trp</sup>	5
tRNA <sup>Arg</sup> /tRNA <sup>Asn</sup>	5
ND4L/tRNA <sup>Thr</sup>	4
ND3/tRNA <sup>Arg</sup>	3
tRNA <sup>Phe</sup> /ND5	1
tRNA <sup>Trp</sup> /tRNA <sup>Tyr</sup>	1
Total	486

greater, 618 bp (Crozier and Crozier, 1993). Thus, comparing the sequenced portion of *M. bicolor* genome with the respective genome portion of *A. mellifera*, we verified that *M. bicolor* presents a more compact arrangement.

The COI-COII intergenic region known to occur in *A. mellifera* was absent in *M. bicolor*. Moreover, indirect PCR evidence indicated that the COI-COII region is absent in at least 16 other Meliponini species (Arias *et al.*, 2006). This intergenic region has been extensively studied in *A.*



**Figure 1** - Linearized mitochondrial genomic map of *M. bicolor* (not in scale). Dots indicate a region not sequenced. Dark boxes indicate tRNA that underwent translocation in comparison to *Apis mellifera* mitochondrial gene order. Numbers above the map indicate the tRNA clusters. I - tRNA<sup>Ile</sup>; A - tRNA<sup>Ala</sup>; K - tRNA<sup>Lys</sup>; M - tRNA<sup>Met</sup>; nd2 - NADH dehydrogenase subunit 2; W - tRNA<sup>Trp</sup>; Y - tRNA<sup>Tyr</sup>; coI - cytochrome c oxidase subunit I; L2 - tRNA<sup>Leu(UUR)</sup>; coII - cytochrome c oxidase subunit II; D - tRNA<sup>Asp</sup>; atp8 - ATP synthase subunit 8; atp6 - ATP synthase subunit 6; coIII - cytochrome c oxidase subunit III; G - tRNA<sup>Gly</sup>; nd3 - NADH dehydrogenase subunit 3; R - tRNA<sup>Arg</sup>; N - tRNA<sup>Asn</sup>; E - tRNA<sup>Glu</sup>; F - tRNA<sup>Phe</sup>; nd5 - NADH dehydrogenase subunit 5; H - tRNA<sup>His</sup>; nd4 - NADH dehydrogenase subunit 4; nd4L - NADH dehydrogenase subunit 4 (light chain); T - tRNA<sup>Thr</sup>; P - tRNA<sup>Pro</sup>; nd6 - NADH dehydrogenase subunit 6; cytB - cytochrome B; S2 - tRNA<sup>Ser(UCN)</sup>; nd1 - NADH dehydrogenase subunit 1; L1 - tRNA<sup>Leu(CUN)</sup>; 16S - rRNA 16S; V - tRNA<sup>Val</sup>; 12S - rRNA 12S.

*mellifera*, and size polymorphism has been reported (from 200 to 650 bp) among subspecies (Garnery *et al.*, 1992, 1995; Franck *et al.*, 1998). It has also been cited as a possible second origin of mtDNA replication and transcription (Cornuet *et al.*, 1991). Our data clearly suggest that this region and possible function is not a shared feature between *A. mellifera* and Meliponini, in a broader sense.

The longest intergenic region found in *M. bicolor* consisted of 173 bp and was located between the tRNA<sup>Met</sup> and ND2 genes. Concerning the sequence similarities between *M. bicolor* and *A. mellifera*, we observed a 46 bp segment within this region that was highly similar (84%) to the non-coding region between COIII and tRNA<sup>Gly</sup> of *A. mellifera*.

The second longest non-coding region of *M. bicolor* mtDNA was located between the ND6 and CytB genes and was 94 bp in length. In *A. mellifera* this non-coding region is 60 bp long. The sequence similarity between these was 61%. The other intergenic regions found in *M. bicolor* were also analyzed, but were too small and showed no significant similarities with any region of the mtDNA of *A. mellifera* or other organisms.

#### Base composition - A+T bias

The adenine+thymine bias was very high in *M. bicolor* mtDNA (86.7%), as has been described in *A. mellifera* (84.9% Crozier and Crozier, 1993). This latter species, as a member of Hymenoptera, has been cited to be the most AT biased insect mitochondrial genome sequenced (Simon *et al.*, 1994). One hypothesis that attempts to explain this bias is that the DNA polymerase could use those bases in a more efficient way during mtDNA replication (Clary and Wolstenholme, 1985). The lower energetic cost to break the A-T links during mtDNA replication and transcription would generate AT bias on organisms that rely on mitochondrial efficiency to keep a high metabolic rate (Xia, 1996). Studies of additional Apidae genomes may indicate whether (and when) this character was fixed by selection for high metabolism in the evolutionary history of this group.

#### Protein-coding genes

The mitochondrial protein-coding genes were analyzed and nucleotide composition, codon usage and size were compared with *A. mellifera*. The initiation codons in *M. bicolor* protein-coding genes were seven ATT (for isoleucine), four ATA, and two ATG (both for methionine). Although the insect mitochondrial genetic code predicts isoleucine for the first codon, it is generally assumed that a special feature on translation changes all mitochondrial initiation codons to methionine on the final amino acid sequence (Wolstenholme, 1992). As in *A. mellifera*, there was no anomalous initiation codon (like *D. yakuba* ATAA; Clary and Wolstenholme, 1985). All *M. bicolor* stop codons are TAA, the standard for the mitochondrial genetic code (Wolstenholme, 1992). There were no incomplete

codons (T or TA), as found in two genes of *A. mellifera* and four of *D. yakuba* (Crozier and Crozier, 1993).

The standard insect mitochondrial genetic code was used to analyze *M. bicolor* mtDNA successfully, since it yielded no stop codons within the gene sequences. The total number of codons (excepting start and stop codons) was 3,643, while the *A. mellifera* genome has 3,686. The codon usage of all *M. bicolor* protein-coding genes was compared with *A. mellifera* (Table 5). It is possible to observe that there is a preferred codon for each amino acid, generally ending with A or T. Interestingly, these codons are not always the complement to their anticodons, particularly when these latter begin with C or G, a feature that is common in insects (Foster *et al.*, 1997). In *A. mellifera*, there are seven codons that are not used at all, and in *M. bicolor* there are even more, 12 non-used codons, all ending with C or G.

The AT bias in codon usage can be expressed in terms of the ratio of "G+C" (Pro, Ala, Arg and Gly) to "A+T" rich codons (Phe, Ile, Met, Tyr, Asn and Lys) (Crozier and Crozier, 1993). That ratio is 0.43 for *D. yakuba*, 0.18 for *A. mellifera* and 0.14 for *M. bicolor*, confirming the extreme AT bias of bee mtDNA. Foster *et al.* (1997) developed a graphical and statistical representation for the nucleotide usage on first and second codon positions, called *square plots*, to analyze AT bias on mitochondrial protein-coding genes. Those graphics do not use the third position because it is too variable and generally does not reflect the amino acid composition. Figure 2 represents the square plot of *M. bicolor*. The AT bias is evident, as most codons are in the first quadrant (two positions occupied by A or T).

Table 6 summarizes the results obtained for all the protein-coding genes of *M. bicolor*, and the comparisons with *A. mellifera*. One of the most intriguing features noticed is the size difference for some genes between the two bees. The *cytB* gene presents the most extreme example, being 102 bp shorter in *M. bicolor*, such difference is concentrated at the amino terminal portion of the *cytB* protein, therefore the reading frame starts 34 codons downstream in reference to *A. mellifera*. Moreover the former species presents a non-coding region of 94 bp preceding the *cytB* gene, with 13 possible initiation codons (nine ATT and four ATA). However none of these give rise to a continuous reading frame. Base substitution or even deletion may explain the absence and presence of the initiation codon at different positions than expected. This difference becomes more striking if one considers that *cytB* is one of the most conserved mitochondrial gene (Simon *et al.*, 1994).

#### Ribosomal RNA genes

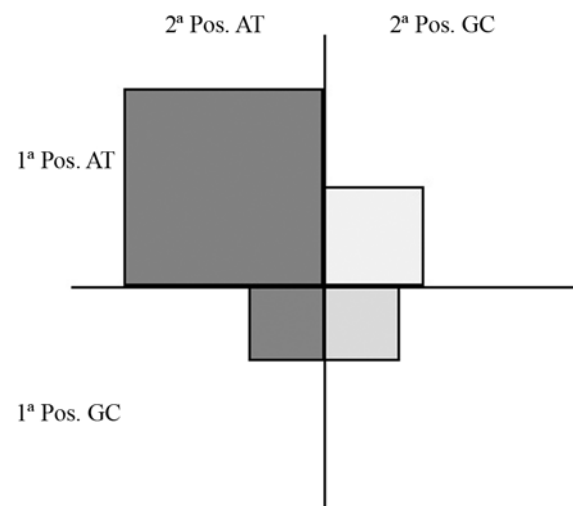
The precise size of ribosomal RNA transcripts are normally difficult to infer from the DNA sequence by itself, so it is assumed that they end on the boundaries of the flanking genes (Boore, 2001). The *M. bicolor* large subunit of ribosomal RNA gene (1rRNA or 16S) was completely sequenced. It has 1,354 bp, 17 bp smaller than the *A.*

**Table 5** - Mitochondrial codons and the absolute number of occurrences on the set of protein-coding genes of *M. bicolor* (Mbi) and *A. mellifera* (Ame). The anticodon of the corresponding tRNA are presented in parenthesis and ? indicates the anticodons not determined for *M. bicolor*.

Amino acid	Codon	Mbi	Ame	Amino acid	Codon	Mbi	Ame	
Ser (?)	AGA	72	81	Gly (ucc)	GGA	82	85	
	AGC	0	2		GGC	0	0	
	AGG	6	2		GGG	1	3	
	AGT	22	18		GGT	29	47	
Ser (uga)	TCA	143	166	Thr (ugu)	ACA	45	72	
	TCC	2	11		ACC	0	5	
	TCG	0	1		ACG	0	1	
	TCT	67	53		ACT	37	53	
Leu (uag)	CTA	12	36	Asp (guc)	GAC	1	5	
	CTC	1	1		GAT	47	52	
	CTG	0	0	Cys (?)	TGC	2	0	
	CTT	13	35		TGT	34	25	
Leu (uaa)	TTA	488	472	Met (cau)	ATA	386	312	
	TTG	15	24		ATG	14	22	
Pro (ugg)	CCA	61	64	Tyr (gua)	TAC	5	10	
	CCC	2	3		TAT	233	209	
	CCG	0	0		Phe (gaa)	TTC	17	26
	CCT	33	36			TTT	423	354
Ala (ugc)	GCA	28	36	Gln (?)	CAA	38	39	
	GCC	1	1		CAG	1	2	
	GCG	0	0		Glu (uuc)	GAA	65	74
	GCT	12	20			GAG	3	5
Arg (ucg)	CGA	28	29	His (gug)	CAC	0	3	
	CGC	0	0		CAT	56	57	
	CGG	1	1		Ile (?)	ATC	21	26
	CGT	8	9			ATT	477	476
Val (uac)	GTA	31	53	Lys (uuu)	AAA	174	152	
	GTC	0	1		AAG	1	8	
	GTG	0	0					
Asn (guu)	AAC	17	11					
	AAT	265	238					
Trp (uca)	TGA	71	78					
	TGG	1	5					

*mellifera* 16S gene (Crozier and Crozier, 1993), and their nucleotide similarity was 81%. The 16S G+C content in *M. bicolor* is 13.2%, while *A. mellifera* is 15.4% and *Drosophila yakuba*, 17%.

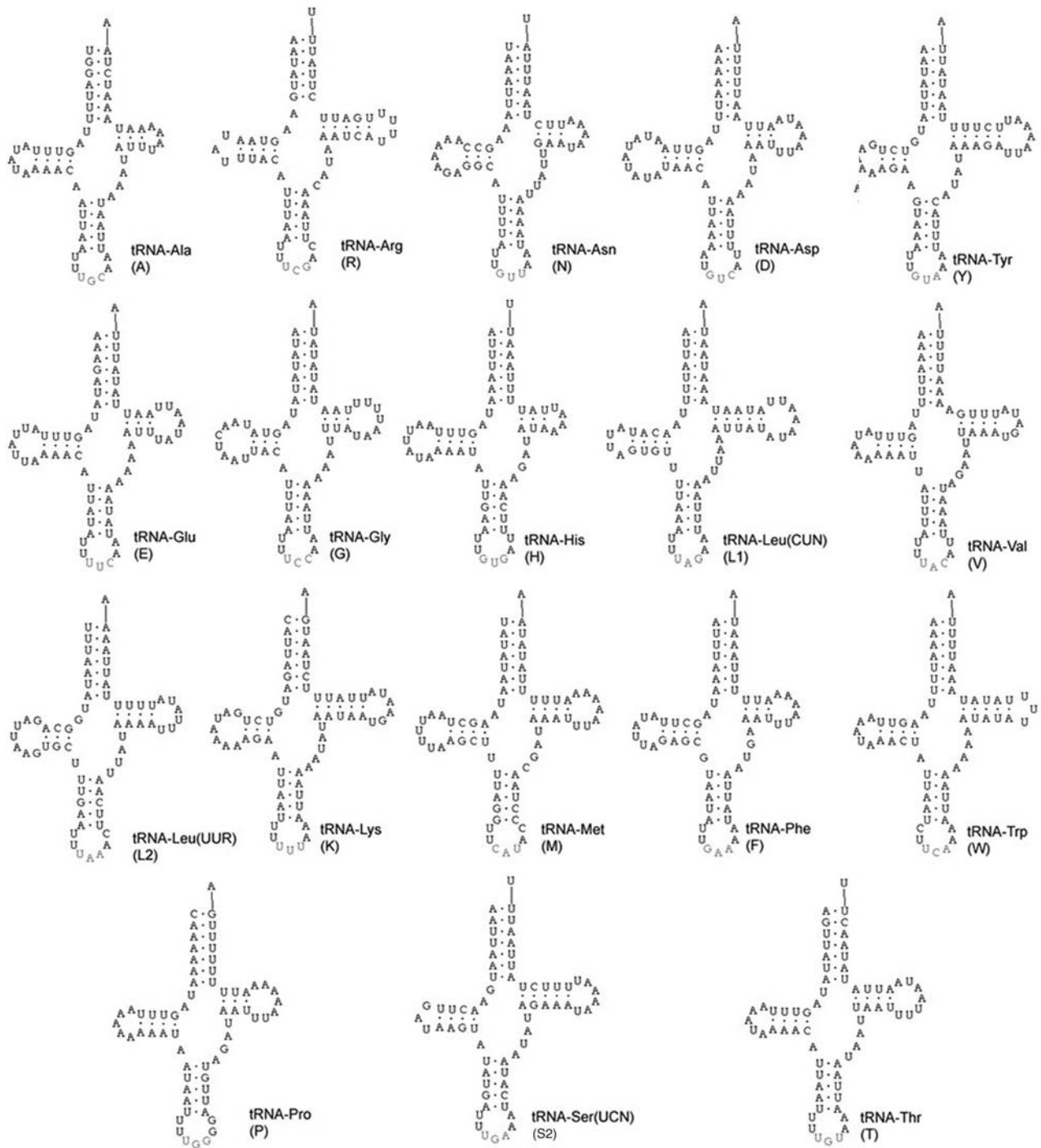
The sequence for the small rRNA subunit (srRNA or 12S) was not obtained completely because it flanks the control region of the mtDNA, which was not amplified. Assuming that this gene has the same length as *A. mellifera*, we sequenced 55% of it (437 bp). The sequence similarity between *A. mellifera* and *M. bicolor* for this sequenced fragment is 79%. As found for 16S, the 12S sequenced stretch presented a higher A+T content (83%) comparing to *A. mellifera* (81%) and *D. yakuba* (79%).



**Figure 2** - Square plot of *M. bicolor* mtDNA. Each quadrant is proportionally dimensioned to the number of codons that are composed by AT or GC on first position (vertical axis) and on second position (horizontal axis).

**Table 6** - Comparisons of protein-coding genes between *M. bicolor* (Mbi) and *A. mellifera* (Ame). Lengths are expressed in basepairs (bp). Start: Start codon; Stop: Stop codon; and Nuc Sim: raw nucleotide similarity between the two bee species. Different start or stop codons are indicated in italic.

Gene	Length Mbi	Length Ame	Start Mbi	Start Ame	Stop Mbi	Stop Ame	Nuc Sim
ATP6	684	681	ATG	ATG	TAA	TAA	79%
ATP8	168	159	ATT	ATT	TAA	TAA	72%
COI	1560	1566	<i>ATT</i>	<i>ATA</i>	TAA	TAA	86%
COII	678	676	ATT	ATT	<i>TAA</i>	<i>T</i>	82%
COIII	780	777	ATG	ATG	TAA	TAA	77%
CytB	1050	1152	<i>ATT</i>	<i>ATG</i>	TAA	TAA	80%
ND1	930	918	<i>ATA</i>	<i>ATT</i>	TAA	TAA	75%
ND2	939	1002	<i>ATA</i>	<i>ATC</i>	TAA	TAA	69%
ND3	354	354	ATA	ATA	TAA	TAA	73%
ND4	1323	1344	<i>ATT</i>	<i>ATA</i>	TAA	TAA	76%
ND4L	279	264	<i>ATA</i>	<i>ATT</i>	TAA	TAA	72%
ND5	1647	1665	ATT	ATT	TAA	TAA	77%
ND6	540	504	ATT	ATT	TAA	TAA	69%



**Figure 3** - Schematic representation of the tRNA secondary structures predicted from the *Melipona bicolor* genome. The anticodon triplet is represented in gray.

The difference in size observed for the 16S gene is quite small, since we have found variations of 102 bp for protein-coding genes (cytB). Size differences are acceptable on rRNA genes more than on protein-coding genes, since there is no need to maintain a frame to read, and only the secondary structure matters to their function (Wolstenholme, 1992). Castro and Downton (2005) aligned the 12S and 16S genes of several insects and found con-

served sequence blocks, indicating that the rRNA secondary structure is also conserved as consequence.

**Difficulties to amplify the A+T-rich region in *M. bicolor***

The A+T-rich region appears generally difficult to amplify in insects, mainly due to tandem repeats, heteroplasmy and great length variation at intra and inter-specific

levels (Zhang and Hewitt, 1997). Also the use of heterospecific primers, designed from flanking tRNA gene sequences, may lead to amplification failure if one of those genes is translocated to another region of the genome (Zhang and Hewitt, 1997).

Our inability to amplify the control region of *M. bicolor* may also be explained by its size. Weinlich *et al.* (2004) estimated it to be about 3,300 bp long, around 2.5 kb longer than in *A. mellifera* (Crozier and Crozier, 1993). Normally such differences are due to partial duplications inside this region, a common feature of insect mtDNA (Simon *et al.*, 1994), which may also make amplification difficult.

### Transfer RNA genes

The tRNA genes were mainly identified by eye, using simple local alignment with their homologues in *A. mellifera* mtDNA. However, when detection was difficult due to low similarity or translocations, these regions were analyzed with specific software (tRNA-Scan, Lowe and Eddy, 1997), which identifies the genes and folds them in typical cloverleaf structures.

From the 22-23 tRNA genes regularly found in animal mitochondrial genomes, 19 were identified on *M. bicolor* mtDNA (Figure 3). The four missing genes were tRNAs for Cys, Gln, Ile and Ser (S1). Although the tRNA<sup>Ile</sup> gene could be identified and positioned on the *M. bicolor* genome, its sequence was not entirely obtained. Analyzing all the *M. bicolor* tRNA sequences as a whole, we have 1,193 bp, with 11.1% G+C and 88.9% A+T. The proportion of A+T for the same genes of *A. mellifera* is slightly less: 87.1%.

The secondary structures of tRNAs of *A. mellifera* and *M. bicolor* were very similar. Nonetheless, a few differences were found, concentrated at the D and TΨC arms, considered the most variable ones (Clary and Wolstenholme, 1985). The anticodon loop always had seven nucleotides, and its arm always had five base pairs, except on *A. mellifera* tRNA<sup>Val</sup> (4 bp). The acceptor arm was also conserved in size (7 bp), except on *M. bicolor* tRNA<sup>Arg</sup> (6 bp). The anticodons were the same for both species.

Eleven tRNA gene clusters (Figure 1) were identified, comprising the 19 tRNA genes sequenced. Nine tRNA genes of *M. bicolor* are on different positions or strands when compared with *A. mellifera*. Two tRNA genes had their position inferred based on their absence in the predicted or sequenced locations. These 11 genes are distributed in only 4 clusters. This number of translocations is higher than those usually found between pairs of Diptera species (OGRE). The molecular and evolutionary implications of this phenomenon on bees are discussed elsewhere (Silvestre *et al.*, 2002; Silvestre and Arias, 2006).

### The KD tRNA cluster in Meliponini

In insects, the junction between the cytochrome oxidase II and ATPase 8 genes normally contains two

tRNA genes, tRNA<sup>Lys</sup> (K) and tRNA<sup>Asp</sup> (D) (*e.g.* Clary and Wolstenholme, 1985). This junction has been called the KD cluster (Dowton and Austin, 1999), as the plesiomorphic organization is considered to be KD (*i.e.* COII-K-D-ATP8); this arrangement has been verified in several members of the Hymenoptera, Diptera and Orthoptera. However, in *M. bicolor* this cluster is represented only by the tRNA<sup>Asp</sup> (D) gene, while *A. mellifera* has the plesiomorphic organization (KD). This observation led us to amplify, sequence and characterize the KD cluster in 14 Meliponini species, representing the large distributional range of the tribe. Species from Brazil, India, Thailand, Australia and Africa were analyzed; all contained only the tRNA<sup>Asp</sup> gene (D). In the *M. bicolor* mitochondrial genome, the tRNA<sup>Lys</sup> gene is located in the first tRNA cluster, near the A+T rich region; this region is considered a hot spot of translocation events (Boore and Brown, 1998). Although we have not sequenced cluster one in the additional meliponine species (to confirm the location of the tRNA<sup>Lys</sup> gene), its absence in the KD cluster is a strong argument that the translocation of this gene occurred very early in the evolutionary history of Meliponini, and seems to be a fixed character of the tribe. Data from other bee families and tribes were obtained (Dowton and Austin, 1999; Silvestre and Arias, unpublished data) and reinforce that this gene rearrangement is a unique feature of Meliponini.

The mitochondrial genome of *M. bicolor* is the first that has been sequenced for stingless bee species. This work provides critical data for future mtDNA analyses of other meliponine species, facilitating the investigation of biological, ecological and evolutionary questions at intra- and inter-specific levels. As several stingless bees are considered endangered, population studies applying molecular tools will be very important in terms of conservation. The sequence obtained here, representing 78% of the total genome, has been available in GenBank since 2003, and has already been used for phylogenetic purposes (Castro and Dowton, 2005) in an attempt to reconstruct the phylogeny of the Holometabola and the position of the Hymenoptera within it. There is an emerging tendency to use complete or nearly complete mtDNA genome sequences for phylogenetic analysis, while this type of data may also provide gene order characters for phylogenetic inferences (Boore and Brown, 1998; Rawlings *et al.*, 2001). It is worthwhile to note the high rate of tRNA gene rearrangement found in the *M. bicolor* mitochondrial genome in comparison to other insects. Studies on wasps have already indicated that the Hymenoptera generally have an accelerated rate of mtDNA gene rearrangement (Dowton and Austin, 1999; Dowton *et al.*, 2002; Dowton *et al.*, 2003). Gene order characters may help resolve the evolution and phylogeny of the bees, particularly the unsolved question about the origin of eusociality among corbiculate bees. The high conservation of the translocation of the tRNA<sup>Lys</sup> gene in Meliponini suggests that it occurred before the great diversification of



Meliponini species and their dispersion over the tropical and southern subtropical areas of the World. The inclusion of more molecular data and analysis of other tRNA clusters may provide further clues for the evolution and biogeography of the Meliponini.

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### Internet Resources

- National Center for Biotechnology Information (NCBI), BLAST searches, <http://www.ncbi.nlm.nih.gov/BLAST>.
- TraceViewer 2.0.1, <http://www.codoncode.com>.
- tRNA-Scan, <http://lowelab.ucsc.edu/tRNAscan-SE/>.

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