Structure and evolution of the mitochondrial DNA complete control region in the *Drosophila subobscura*

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Abstract

The complete A + T-rich region of mitochondrial DNA (mtDNA) has been cloned and sequenced in the species of the *Drosophila subobscura* subgroup *D. subobscura*, *D. madeirensis* and *D. guanche*. Comparative analysis of these sequences with others already published has identified new sequence motifs that are conserved in *Drosophila* and other insects. A putative bi-directional promoter and a stop signal are proposed to be involved in the primary mtDNA strand replication of *Drosophila*. This region strongly resolves relationships of the species included in a phylogenetic analysis, both for closely related species and also at deeper phylogenetic levels when only the left and central domains are taken into account.

Keywords: *Drosophila*, control region, promoter, phylogeny.

Introduction

The Control Region (CR) is the only large non-coding region in the mitochondrial DNA (mtDNA). This region has been the object of numerous functional studies in vertebrates, which have identified the transcription initiation sites for each strand (Chang & Clayton, 1984), and the main origin of replication (Clayton, 1982). In invertebrates it has been less studied, although in the crustacean *Artemia franciscana* a main and several potential transcription initiation sites have been located in its non-coding region

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(Carrodeguas & Vallejo, 1997). In Drosophila, in which the CR is also known as the A + T-rich region, the only regulatory feature unequivocally localized is the unidirectional origin of replication that was mapped by electron microscopy in several species (Goddard & Wolstenholme, 1980). In addition, a secondary stem and loop structure has been proposed as the initiation site of the second strand synthesis (Clary & Wolstenholme, 1987) by analogy with mammals (Martens & Clayton, 1979). Comparative studies in vertebrates and invertebrates have subdivided the CR into three domains with different levels of variability. A left domain exists adjacent to the tRNA^{lle} gene in Drosophila, the central domain is the most conserved in all taxa, and the third domain, usually highly variable, is 5' upstream of the small rRNA gene in Drosophila. In mammals, Doda et al. (1981) have described termination-associated sequences of the H-strand replication in the left domain. No function has yet been assigned to the left domain in Drosophila. The central subregion shows extended nucleotide sequence similarity through several taxonomic levels in vertebrates and in Drosophila. Finally, in spite of the high variability of the right domain, it is there where the most conserved sequence blocks in vertebrates are found, which are implicated in several aspects of the mtDNA replication. Although a general model of the CR functions has been proposed for vertebrates (Shadel & Clayton, 1997), it seems that it is not extensible to invertebrates. In Drosophila, the complete CR has only been sequenced in D. virilis (Clary & Wolstenholme, 1987), in some species that have much shorter A + T-rich regions (Clary & Wolstenholme, 1985; Monnerot et al., 1990), and in D. melanogaster which has a 4.6 kb long A + T-rich region (Lewis et al., 1994).

Here, we describe the complete CR sequence of the related species *D. subobscura*, *D. madeirensis* and *D. guanche* that compose the *subobscura* subgroup of the *obscura* group. We compare the phylogenetic estimate of relationships derived from the CR with that previously suggested based on other mitochondrial and nuclear DNA sequences. We also determine structural patterns in the CR, both within the *subobscura* group and with other *Drosophila*.

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SubR SubL mad gua	GCAATTCATT Region A AATAGTGAGATTCTCTCTCTAT-TTTTTTTTTTTTTTTTT	104 104 105 104
SubR	Region B TTAATTAATAATAGAATTTAGTAATTTTTATTTATTATAAAATTATT	207
SubL mad gua		207 208 207
	tatataaatatataaatatataaatatataaata llllllllll	
SubR SubL mad gua	TATATATAAATATATAAATATATAATAATTTAAAAATAATTATTAAAA	312 312 313 312
SubR SubL mad gua	TGTCACCATTTTTGGATT GCGAAATTTTTTATTTAAATTGTTAACTATATAAAGATATATTAAACATTTATATATA	417 417 418 417
SubR SubL mad gua	Region C ATATAAAAAAAAAAAAA TAGTAATTTTTTAATATATATATATAAAATTACTATTAATTTATTTAAATT-ATTTAATTATATATA	515 515 517 515
SubR SubL mad gua	ATATAAATATAAATTAT - TATTTTGATATTATTAATATTAATTAATTAATTAATTAA	618 618 619 615
SubR SubL mad gua	ТТТАТТАТТААТАААААТТ-АТСАТТТАТТАА-АААТТАТТАТТАТТАТТААТАААТТАТАААТАААТАААТАААА	721 721 720 718
	atatttatttata aattag ^g g g tataaataaa-at _{tt} ata _t tc _a g	
SubR SubL mad gua	TAAAATAAAGAATTTTTTATAAAATA-AAAATTTA-TATTCAGGGGGATTAT-AGATATTTATTATAAAAATTAAAATTTATAAATTTAT-ATTTAATTTT	821 821 817 813
SubR SubL mad gua	TATCAGATTTAGTAATTAAAATTATTATATTTTTGTGTATTTTAATTATT	926 926 922 916
SubR SubL	TATTAAAAAATTACTATTTTATTTTTAAAGTTTTATTT 964 	

Figure 1. Alignment of the nucleotide sequences analysed. A *dot* indicates a nucleotide that is the same as that in *D. subobscura* from Raíces. A *dash* indicates a nucleotide that is absent. A *letter* indicates a substitution. *Brackets* indicate the limits of the three putative regions. *Dashed brackets* indicate the limits of the sequence that corresponds to the stem-loop structure. The palindromic and putative stop sequences are underlined. The most stable secondary structures are indicated above the stem-loop, the palindromic and the putative stop sequences.

Results

Structure of the A + T-rich region

Length of the CR in the subobscura subgroup ranges from 930 nt in D. guanche to 940 nt in D. subobscura. The overall A + T content for the entire CR in the D. obscura group is 93%, which is typical of this region in Drosophila. Alignment of these sequences gives a total length of 963 nt (Fig. 1). A screening of nucleotide variation along the sequence using a 20 nt window with the MEGA program (Kumar et al., 1993) allows the identification of three regions of different variability. The first, A (bases 11-157, Fig. 1) is immediately adjacent to the tRNA^{lle} gene and has the highest nucleotide diversity (0.230 ± 0.058) ; the central, B, bases 158-437, is the most conserved (0.021 ± 0.007) ; and the third, C, bases 439-973, upstream from the 12S rRNA gene, is more variable (0.150 ± 0.041) than B but less than A. The rate of nucleotide substitution for the regulatory region as a total in the subobscura subgroup (0.116 ± 0.03) is four times faster than that of mitochondrial coding regions (0.029 ± 0.006) for the same species (Barrio *et al.*, 1994) but the conserved B domain (0.021 \pm 0.007) is at the same level.

Phylogenetic analysis

For phylogenetic comparisons we included from GenBank sequences from *D. obscura*, *D. ambigua*, *D. yakuba*, *D.*

teissieri, D. melanogaster, D. simulans, D. mauritania and D. virilis. It was not possible to align region C, which varies in length from approximately 550 bp in the suboscura group to 3.5 kb in D. melanogaster. Furthermore this region is not reported for some of the taxa. However 434 bp (regions A and B) were included in the analysis for the twelve taxa. Using Modeltest (Posada & Crandall, 1998) under the Akaike information criteria we concluded that the TrN model (equal transversion ratios, transition ratios A/G 4.1, C/T 2.3 relative to transversions), with a gamma distributed rate heterogeneity model (four rate categories, G = 0.85; Yang, 1994), was the most appropriate model of evolution for this data. A ten replicate heuristic search with this model produced a single tree with a maximum likelihood score of -1757 (Fig. 2).

Discussion

Putative function of conserved elements in the A + T-rich region

Previously the only conserved element known in region A of *Drosophila* was a thymidylate stretch with a mean of 21 ± 2 nts near the tRNA^{lle} gene. This run of Ts is also present in the *subobscura* subgroup species with similar number and position (Fig. 1). Due to its localization, this stretch might be expected to form part of a promoter for transcription in the direction from the A + T-rich region to



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the tRNA^{lle} gene (Clary & Wolstenholme, 1987). The central domain B contains the largest and most conserved region among all the Drosophila species studied (Fig. 1). Clary & Wolstenholme (1987) identified in the centre of this region a possible stem-and-loop structure, which was suggested as a possible secondary origin of mtDNA replication in Drosophila. We detected a highly conserved palindromic structure localized within the B region at the left of this putative stem-loop motif, which is not only present in all the Drosophila species studied (underlined in Fig. 1) but also in the same relative position in other insects such as Anopheles and Locusta. It is one of the most favourable secondary structures detected in the region and its mean free energy is higher (DG = -14) than those described previously. The right border of the B region ends in a very conserved run of As, which in the subobscura species has a mean length of 14 ± 1 nt, which is within the range of all the Drosophila species examined (15 ± 4 nts). Due to its position adjacent to the innermost conserved type II repeat in D. melanogaster, a role in replication has been suggested for this A-stretch (Lewis et al., 1994). The C region has little homology even within the Drosophila species studied. Alignment of this region is only possible among closely related species such as those of the subobscura subgroup in this paper (Fig. 1) or among those of the *melanogaster* subgroup (Inohira et al., 1997). No conserved sequence motifs have been described for this region. However, we have detected in it a well conserved block that is not only present in the subobscura subgroup species and in all the Drosophila sequenced for this region, but in other distantly related insects. The motif is a stretch of four or five Gs. In addition, although the primary sequences of the flanking regions are different in distantly related species, a stem-loopstructure with the Gs displayed in the loop is possible in all of them (underlined in Fig. 1). In the fast evolving region C, the presence of the G stretch stands out, not only for its maintenance across unrelated taxa but also for the oddity that such a motif exists in a region that is 95% A + T. From the primary sequence, secondary structure and position 3' downstream of the primary origin of replication, this G motif might be associated with the stop of the primary strand replication in Drosophila and other invertebrates.

Mitochondrial DNA promoter sequences are generally not well conserved except for their A + T rich composition (Jaehning, 1993). We performed a search for possible promoters using the NNPP program (Reese *et al.*, 1996), concentrating on putative promoters that are localized in similar positions and orientation in different species. A pair of symmetric promoters, one in each strand, appear localized in region B in the conserved block upstream of the stem-loop motif, with the consensus sequence AYRT TATA. This sequence also appears in phylogenetically distinct organisms such as yeast (Christianson & Rabinowitz, 1983) and chickens (L'Abbé *et al.*, 1991).

Phylogenetic value of the A + T-rich region

Within the suboscura group D. madeirensis is most closely related to D. subobscura (99% bootstrap support) and D. guanche is the sister group to this pair (95% support). This is concordant with topologies derived from other mitochondrial and nuclear gene regions (O'Grady, 1999). However, despite the relatively short segment used, bootstrap support for relationships is much higher than obtained with other mitochondrial genes even those for which longer sequences were included (O'Grady, 1999). Therefore while the C region of the control region may not be phylogenetically useful the other parts are very effective at recovering the mtDNA tree. Caution should still be exercised when comparing this to the species tree however, because some of these taxa, for example D. simulans and D. mauritana, have been shown to have introgressed mtDNA (Ballard, 2000). Clearly in this case the gene tree will not necessary correspond to the species tree.

Experimental procedures

Laboratory methods

Two specimens of each species were sequenced for the entire A + T-rich region. For the widespread D. subobscura species one strain (subR) was from Tenerife (Canary Islands) and the other (subL) from Lisbon (Portugal). For the endemic species D. guanche (gua) and D. madeirensis (mad) the specimens were from the Tenerife island and Madeira island, respectively. Total DNA was extracted from individual adult flies by the CTAB method (Towner, 1991) with minor modifications. The entire A + T-rich region was amplified in two overlapping fragments: Fragment I, closest to the tRNA^{lle} (450-451 bp), using primers R1 (5'-CCTAT-CAAggTAACCCTTTTTATCAg-3') and R2 (5'-gTgTATACTAAgT-TCTAAAT TAATAg-3' - Monforte et al., 1993). Fragment II, closest to the small rRNA was amplified using primer R10 (5'-TgTAAC-CATTTTTggATTgCgA-3') designed from our sequences, and R8 (5'-AACTAATAACAAATTTTTAAgCCA-3', Clary & Wolstenholme, 1987). PCR amplifications were performed in a mix reaction containing 0.1 mm of 7-deaza dGTP and dCTP, and 0.3 mm of dATP and dTTP. In the cycle profile reduced extension temperatures are typically used to amplify the A + T-rich region (Xin-Zhuan et al., 1996) -- 20 s at 90 °C, 1 min at 48 °C and 1 min at 68 °C for thirtyfive cycles. In order to read accurately the sequences after the stretch of Ts, found at the beginning of fragment I, we proceeded to clone these fragments in the plasmid pBS II KS (+/-). In the presence of dTTPs alone, Tag DNA polymerase was used to add a single T to the blunt ends of the EcoRV digested vector in order to clone efficiently the PCR products (Ausubel et al., 1995). The ligation was performed with the Ready to Go T4 ligase kit (Pharmacia) and the ligate was used to transform the XL1-blue E. coli strain (Stratagene). Nucleotide sequences were obtained by the double-stranded dideoxy terminator procedure with the Promega fmol DNA sequencing system, using primers isotopically (γ-³²P) ATP end labelled. Because of the high level of A and T in the sequences, modified nucleotide mix was used, containing 30 mM of dATP and dTTP, and 3 mM of dCTP and 7-deaza dGTP. The nucleotide mix d/ddG was 9 mM ddGTP, d/ddA was 900 mM ddATP,

d/ddT was 1500 mM ddTTP and d/ddC was 45 mM ddCTP. The sequencing reactions were performed using the cycle profile: 15 s at 92 °C, 15 s at 52–55 °C and 15 s at 65 °C for forty-five cycles. In addition to the PCR amplification primers, R11 (5'-ATATAgTTgA-TAATTTATAAACA-3') and R7 (5'-AATAAATATTTATAATCCCCCT-gAAT-3') were used to sequence completely fragments I and II, respectively. Clones were sequenced with the M13 universal primers.

Structural and phylogenetic sequence analysis

The new nucleotide sequences reported here are available on GenBank, accession numbers AJ132899-AJ132902. Previously published sequences from other Drosophila for the same region were imported from GenBank. Multiple sequence alignments were performed with CLUSTAL X (Thompson et al., 1997) and later maximized for sequence similarity by visual inspection. The MFOLD program (Mathews et al., 1999) was employed to identify potential secondary structures. Putative promoters were determined using the program NNPP (Reese et al., 1996). Sequences were phylogenetically analysed using PAUP* (Swofford, 2001). When estimating phylogenetic relationships among sequences, one assumes a model of evolution regardless of the optimality criteria employed. We used the approach suggested by Huelsenbeck & Crandall (1997) to test fifty-six alternative models of evolution, employing PAUP* and Modeltest (Posada & Crandall, 1998), outlined in detail in Harris & Crandall, 2000). Once a model of evolution was chosen, it was used to estimate a tree using maximum likelihood (Felsenstein, 1981). Confidence in resulting nodes was assessed using the bootstrap technique (Felsenstein, 1985) with 1000 replicates.

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