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# A compact gene cluster in *Drosophila*: the unrelated *Cs* gene is compressed between duplicated *amd* and *Ddc*

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### **Abstract**

*Cs*, a gene with unknown function, and *amd* and *Ddc*, which encode decarboxylases, are among the most closely spaced genes in *D. melanogaster*. Untranslated 3' ends of the convergently transcribed genes *Cs* and *Ddc* are known to overlap by 88 bp. A number of questions arise about the organization of this tightly-packed gene region and about the evolution and function of the *Cs* gene. We have now investigated this three-gene cluster in *Scaptodrosophila lebanonensis* (which diverged from *D. melanogaster* 60–65 MYA), as well as in *D. melanogaster* and *D. simulans*. Gene order and direction of transcription is the same in all three species. The *Cs* gene codes, in *Scaptodrosophila*, for a polypeptide of 544 amino acids; in *D. melanogaster*, it consists of 504 amino acids, which is twice as long as previously suggested, which makes the gene density even more spectacular. The *Cs* sequences exhibit higher number of non-synonymous substitutions between species, higher ratios of non-synonymous to synonymous substitutions, and lower codon usage bias than other genes, suggesting that *Cs* is less functionally constrained than the other genes. This is consistent with the failure of inducing phenotypic mutations in *D. melanogaster*. The function of *Cs* remains to be identified, but a high degree of similarity indicates that it is homologous to genes coding for a corticosteroid-binding protein in yeast and a polyamine oxidase in maize. © 1999 Elsevier Science B.V. All rights reserved.

*Keywords:* Decarboxylases; *D. melanogaster*; Gene cluster; Gene duplication

the left arm of the second chromosome, includes 18 by gene duplication (Eveleth and Marsh, 1986). An identified genes plus three transcription units for which enigmatic gene, called *Cs*, lies between *amd* and *Ddc* no detectable phenotypic mutations are known (Maroni, (Eveleth and Marsh, 1987). All three genes are among no detectable phenotypic mutations are known (Maroni, (Eveleth and Marsh, 1987). All three genes are among 1993; Wright, 1996; Stathakis et al., 1995). Most of the most closely spaced genes in *D. melanogaster*, and 1993; Wright, 1996; Stathakis et al., 1995). Most of the the most closely spaced genes in *D. melanogaster*, and genes are densely clustered in two subclusters. Many the 3' ends of the *Ddc* and *Cs* genes actually overlap genes are densely clustered in two subclusters. Many the 3' ends of the *Ddc* and *Cs* genes actually overlap by genes in the cluster are functionally related in that they as by (Spencer et al. 1986a: Stathakis et al. 1995 genes in the cluster are functionally related in that they 88 bp (Spencer et al., 1986a; Stathakis et al., 1995). In are involved in the catecholamine metabolism.

Two genes from the proximal subcluster, *Ddc* and<br>amd, have been well studied, with about 90 phenotypic<br>isolated mutations (Wright, 1996). Four genes from the<br>proximal subcluster, including *Ddc* and amd, have been<br>*Ddc* h

**1. Introduction** sequenced in *D. melanogaster* (Eveleth et al., 1986; Marsh et al., 1986). The coding regions of these two The *Ddc* gene cluster in *D. melanogaster*, located on genes are highly similar and are thought to have arisen are involved in the catecholamine metabolism.<br>Two genes from the proximal subcluster, *Ddc* and  $\frac{1}{2}$  known for *Cs*. The product of the *Cs* gene is not known

from mammals to insects, including *D. melanogaster*. Until now, the *amd* has been studied only in *D. melano-* Abbreviations: *amd*, <sup>a</sup>*-methyl dopa sensitive* gene encoding decarbox-*Ddc*, gene encoding Dopa decarboxylase (DDC, EC 4.1.1.26); ENC,<br>
effective number of codons; Myr, million years; MYA, million years<br>
ago; PCR, polymerase chain reaction; PLP, pyridoxal 5'-phosphate. (Jackson, 1990), no ge \* Corresponding author. Fax: <sup>+</sup>1-949-824-2474. similar to *Cs*. The origin of *Cs* is unknown. Its position *E-mail address:* antatare@uci.edu (Andrey Tatarenkov) between *amd* and *Ddc* could be a consequence of the

ylase related enzyme (product unknown); bp, base pair(s); BLAST, *gaster*, and the *Cs* gene is only known to occur in *D.* basic local alignment search tool; *Cs*, a gene with unknown function; *melanogaster*. While *Ddc* and *amd* are members of a

original *amd*–*Ddc* duplication; or it may have been *2.2. DNA preparation and sequencing* inserted there at a later time (Eveleth and Marsh, 1987).

Thus, a number of questions arise about *Cs* and its Total genomic DNA was obtained using the phenol– function and location within a developmentally impor- chloroform extraction procedure described by Palumbi tant gene cluster. The first question is whether the *Cs* is et al. (1991). To design amplification primers, we compresent between *amd* and *Ddc* in other species as well, pared published sequences of *Ddc* from the moth as it is in *D. melanogaster*. Second, the compactness of *Manduca sexta* (GenBank accession number U03909), the *Cs*, *amd*, and *Ddc* cluster in *D. melanogaster* is the mosquito *Aedes aegypti* (U27581), and *D. melano*unusual, and it is of interest to find out whether this is *gaster* ( X04661), as well as the *amd* from *D. melanogas*a result of recent events, or, rather, whether such com- *ter* ( X04695). *Ddc* and *amd* in *D. melanogaster* are quite pactness is old, perhaps tracing back to the time of the similar to each other in sequence but have different *amd*–*Ddc* duplication. One more question concerns the orientation. We selected segments of the aligned functional role of the *Cs*. As Li (1997, p. 185) has sequences, that had high similarity but also specific pointed out, it is well known ''that the stronger the substitutions in the *amd* sequence when compared functional constraints on a macromolecule, the slower with *Ddc* sequences. The two primers (forward the rate of evolution". Thus, if the *Cs* has a less vital 5′-GAYATYGARCGNGTSATCATGCCKGG-3′, and function for the organism than *amd* and *Ddc*, it is reverse 5′-GAYATYAGYCGNGTSATCAAGCCKexpected that its evolution be faster than that of the GG-3∞) encompass large parts of *Ddc* and *amd* as well two neighboring genes. Moreover, investigating the as the interval between them (Fig. 1). A region of about pattern of substitutions could help to ascertain whether 5.8 kb was obtained in several species of Drosophilidae. the  $Cs$  is a protein encoding gene, which has been PCR reactions were performed in a  $100 \mu$  volume

whole *amd–Cs–Ddc* cluster, in the Drosophilid ward and reverse primers,  $0.2 \text{ mM dNTP}$ , and  $3 \mu$  of *Scaptodrosophila lebanonensis*, from a genus closely genomic DNA. The cycling parameters for the amplifirelated to *Drosophila*. We have also sequenced in *D.* cation were an initial denaturation at 95°C for 5 min *melanogaster Ddc* and the coding region of *Cs* in order and 31 cycles of the following: denaturation for 30 s at to resolve inconsistencies arising from previous pub- 95°C, annealing for 1 min at  $60^{\circ}$ C, and extension for lished sequences. Finally, we have also sequenced most 5 min at  $72^{\circ}$ C for the first cycle and an extra 3 s for of the three-gene region in *D. simulans* for the purpose every subsequent cycle; after 31 cycles the reaction was of confirming inferences about *D. melanogaster*. additionally kept at 72°C for 7 min to complete

Comparison between the *Cs* genes of *S. lebanonensis* extension. and *D. melanogaster* shows high sequence similarity The PCR product of *S. lebanonensis* was purified between them, comparable with the similarity observed with Wizard PCR preps DNA purification system for the neighboring *Ddc* and *amd* genes. Moreover, the (Promega Corporation), and cloned using the TA clonregions of high similarity in the nucleotide and putative ing kit (Invitrogen, San Diego, CA). DNA sequencing amino acid sequences extend much beyond the coding was partly done by the dideoxy chain-termination techregion previously suggested for *Cs* (Eveleth and Marsh, nique with Sequenase Version 2.0 T7 DNA polymerase 1987). It follows that the three genes are even more (Amersham Life Sciences Inc., USA) using <sup>35</sup>S-labeled tightly packed than had been previously thought for *D.* dATP, and partly with an ABI model 373 autosequencer *melanogaster*, and that they are partially overlapping. using Dye Terminator Ready Reaction Kit in accordance

questioned (Eveleth and Marsh, 1987). of the ExTAKARA buffer containing 2.5 U of We have sequenced the *Cs* gene, as well as the ExTAKARA Taq polymerase,  $0.5 \mu M$  each of the for-

with the manufacture protocol (Perkin Elmer) (see Fig. 1). We employed a successive approach for sequencing the region, so that new sequencing primers were designed based on the sequence obtained with previous **2. Materials and methods primers.** Both strands were completely sequenced with 34 primers.

2.1. Species Sequences of the *Cs* gene in both *D. melanogaster* and *D. simulans* were obtained by direct sequencing of Isofemale lines of *Drosophila melanogaster*, *D.* purified PCR products with an ABI model 377 auto*simulans*, and the closely related Drosophilid sequencer using the Dye Terminator Ready Reaction *Scaptodrosophila lebanonensis* were studied. *D. melano-* Kit in accordance with the manufacturer's protocol *gaster* and *D. simulans* were collected by one of us (Perkin Elmer). Partial sequences of *Ddc* in *D. melano-* (FJA) in St. Lucia, West Indies, in 1995. The strain of *gaster* and *D. simulans* were obtained from separately *S. lebanonensis* is from the National Drosophila Species constructed clones of these species. The sequences of Stock Center in Bowling Green, Ohio. these clones overlap considerably with the PCR frag-



Fig. 1. Structure, gene arrangement, and direction of transcription of a genomic DNA segment comprising the genes *amd*, *Cs*, and *Ddc* in *Scaptodrosophila lebanonensis* and *Drosophila melanogaster*. Thick arrows adjacent to gene symbols indicate direction of transcription from 5∞ to 3′. Boxes indicate protein coding regions: thick lines connecting them represent introns; thin lines represent the non-coding regions. Dotted lines connect the *Cs* regions of high similarity between the two species. The two thick lines in the lower part indicate regions that we have sequenced in *D. melanogaster* and *D. simulans*; the rest of the melanogaster sequence is from Marsh et al. (1986) and Eveleth and Marsh (1987). The gene structure and arrangement are the same in *D. simulans* as in *D. melanogaster*.

ments. Partial sequence of *amd* in *D. simulans* was *ter* ( U18307) and *D. pseudoobscura* ( X16337). Codonobtained from yet another clone, which is encompassed use bias was assessed by estimating ENC, the 'effective by the PCR fragment. The number of codons' (Wright, 1990). Higher values of  $\alpha$ 

GenBank database, accession numbers AF091327, of substitutions along amino acid sequences was tested AF091328, AF091329, AF121109. with the unmodified variance test of Goss and Lewontin

grams of the Fragment Assembly module of the GCG BLAST at http://www.ncbi.nlm.nih.gov/. package ( Wisconsin Package Version 9.1). Various GCG programs were also used for alignment and translation. Inference about coding regions was primar- **3. Results** ily obtained by comparison of the *S. lebanonensis* and *D. melanogaster* sequences seeking regions of high sim- A DNA fragment of approximately 5.8 kb resulted ilarity. Additionally, the programs GENIE (Reese et al., from PCR amplification in several drosophilid species, 1997) and FGENED (Solovyev et al., 1994) were used *Scaptodrosophila lebanonensis*, *D. melanogaster*, *D. sim*for predicting putative exons. Analysis of codon prefer- *ulans*, *D. immigrans*, *D. mimica*, *D. (Scaptomyza)* ence was performed with the CODONPREFERENCE *palmae*, and *D. (Samoaia) leonensis*. The gene organizaprogram of the GCG package which implements the tion of the amplified region in *D. melanogaster* and *S.* method of Gribskov et al. (1984). A Fourier transform *lebanonensis* is outlined in Fig. 1. analysis was performed using the Fast Fourier We searched the region between the stop codons of Transform of the computer program Origin (version *Ddc* and *amd* in *S. lebanonensis*, presumably correspond-4.10, Microcal Software, Inc.). This method unveils ing to the *Cs* gene, seeking segments similar with the periodicity patterns along binary strings. Such strings sequence of *Cs* in *D. melanogaster* ( X05991). We found were created by using a 1 at each substituted position, an extended region, about 1.5 kb with high similarity and a 0 at identical positions. In addition to the aligned (71%) to the sequence of *Cs* in *D. melanogaster* (Figs. 1 coding regions of *amd*, *Ddc*, and *Cs* of *D. melanogaster*, and 2). Unexpectedly, the region of similarity extends *D. simulans*, and *S. lebanonesis*, we also used for illustra- more than 400 bp beyond the previously suggested *Cs* tive purposes *hsr-omega* exons 1 and 2 of *D. melanogas-* stop codon in *D. melanogaster* (Eveleth and Marsh,

The sequences reported here have been deposited in ENC correspond to lower codon-use bias. Heterogeneity (1996). The analysis was kindly conducted by R.C. *2.3. Alignment and analysis* Lewontin. Rates of substitution at synonymous and non-synonymous sites were calculated by the method of The sequences were edited and assembled using pro- Li (1993). We searched GenBank sequences with the

MEI STM LEB T TTT AAA CTG GCC AGC AGC AGC GCC AGC TTA TAC AAT GCA CGG GTT CTA CAG --- --- --- --- --- --- --- --- 153 MET. STM LEB MEL GCG GAT AAC ATC GGC GAC AAG CAA CGC AGT CCA GAT CTG GAG GCG GCG CGC CAA AAT ACC CAG ATA GTG GTC GTG GGC 231 MEL GCA GGA CTC GCC GGT CTC TCG GCG GCC CAG CAC CTC TTG TCG CAC GGC TTT CGG CGC ACT GTG ATC CTG GAG GCC ACA 309 MEL GAT CGT TAT GGC GGC AGG ATT AAC ACC CAG CGC TTT GGT GAC ACC TAC TGT GAA CTA GGC GCC AAG TGG GTA --- --- 387 MEL AAG ATC GAT GGA TCG CAG GAC TCA ATG TAT GAA CTG CTA CGC AAC ACG GAA GGC TTG GGG AAG CAG ATA AAG CAG CCG 465 MEL GAT CGG GCC ACC TAT CTT CAG --- GAT GGA AGC CGC ATC AAT CCA GCC ATG GTC GAG CTT ATC GAC ACG CTA TTT CGG 543 MEL CAG CTT TGC QGA GGC TTC AAG GTC TCC GAA CGA GTT AAA ACG GGT GGT GAC CTG CAC TCG CTG GAC AAT GTC ATG AAC 621 MEL TAC TTT AGA ACA GAA AGC GAT CGC ATC ATT GGC GTC TCC TTC CAG CAT CCT AAG GAT CAA CTG GCG GCA CGC GAG ATC 699 MEL TTC CAA TCG CTG TTC AAG GAG TTC GGC AGC ATC TTG GGA TGC TGC CTG GAG TAC GTG AAC ATC GAA CAC ATA ACC AAG 777 MEL TGT CCA GTG CAG CAG GAA CAG CGC CCG CGT TAT GTG CCC ACT GGT CTA GAT AAT GTA GTG GAC GAT CTC ATT CAG AAC 855 MEL --- ACA CCA GCG CCG ATG --- --- AAA AGT GTG GGT TGC CTG GAT GGC AGT CTT TAC AAC GCC GAT CAC ATA ATA TGC 1011 MEL ACC CTG CCG CTG GGC GTG CTC AAA AGC TTT GCG GCC GTT CTG TTT CGA CCC ACG CTG CCG CTG GAC AAG ATG CTG GCT 1089 MEL ATA CGC AAC CTC GGC TTT GGC AAT CCC CTC AAG ATA TAT CTC TCC TAC AAG AAG CCC ATT GGG CGT TGG CTA AAG GGA 1167 MEL ACC CTC CGG CCA CTG GGA ACG CTT CTG --- --- --- AAT CCT --- --- TCC GTG GAG CAG CAA CCC GAA CGC AAC TGG 1245 MEL ACG CAG CAG GTC GTG GAG ATC AGC CAG GTG CCC AGC AGT CAG CAT GTG CTG GAG GTG CAT GTC GGT GGC GGA TAC TAC 1323 MEL CTG GTG CCG TAC CCA CAG GAA CTG CTG CGT TCC AAC TGG AGT ACG TCG GCC TGC TAC CTC GGC GGT CGT CCT TAC TTC 1479 MEL GGG GAT GCA ACC TCG TTG AGA AGC TTT GGA ACC ATT GAT GCC GCC AGG TCC AGT GGC ATC CGA GAA GCC CAA CGC ATC 1635 

Fig. 2. Alignment of the *Cs* coding region between *D. melanogaster* (MEL), *D. simulans* (SIM), and *S. lebanonensis* (LEB). The intron in *D. melanogaster* and *D. simulans* is shown in lowercase letters at the top of the figure; the proposed initiator ATGs are underlined; the stop codons are in bold. A region of uncertain alignment is overscored with a double dotted line (top). Dots indicate nucleotides identical to *D. melanogaster*; hyphens indicate gaps. Discrepancies between our *Cs* sequence and the sequence of *D. melanogaster* of Eveleth and Marsh (1987) are shown with rectangles to indicate nucleotides missing in their sequence, and arrows to indicate locations at which they show excessive number of nucleotides.

1987). Moreover, the *S. lebanonensis* sequence has high of the first exon merge with the remaining 1507 bp of similarity to a segment upstream of the largest ORF the long ORF that we have found in *D. melanogaster*. previously identified (Eveleth and Marsh, 1987) in the FGENED suggests that the coding region of *Cs* in *S. D. melanogaster Cs* gene. This whole 1.5 kb region is an *lebanonensis* consists of just a single exon, which starts uninterrupted open reading frame (ORF) in *S. lebano-* 21 codons upstream of the region where similarity *nensis*. While the *S. lebanonensis* and *D. melanogaster* between *D. melanogaster* and *S. lebanonensis* can be sequences are highly similar at the nucleotide level along detected. GENIE yields the same start and stop codons the whole 1.5 kb region, the corresponding peptide as FGENED. However, GENIE indicates the presence sequences are similar only in a few stretches, which are in *Scaptodrosophila* of a short intron (positions 587– interrupted by stretches that cannot be aligned. This 718 in Fig. 2). We rather assume that this is a coding appears to be a consequence of shifts in reading frame segment given that it is highly similar to the sequences due to indels in the published sequence of *D. melanogas-* of *D. melanogaster* and *D. simulans* along the segment's *ter* (Eveleth and Marsh, 1987) compared with *S.* whole length at both the nucleotide and the amino acid *lebanonensis*. level. It is also possible that the *Cs* in *D. melanogaster*

gene and adjacent regions in *D. melanogaster*, as well as that an intron (62 bp) may have arisen due to mutations in the closely related *D. simulans*. Our *Cs* sequence of that have disrupted the beginning of the coding *D. melanogaster* differs from the published sequence by sequence. This would explain the somewhat unusual the occurrence of nine indels, as predicted by the align- position of the intron, after an exon of only three ment of the previously published sequence with the *Cs* codons. It is also possible, but seems less likely, that an sequence of *S. lebanonensis* (see Fig. 2). intron in the ancestral species may have become a coding

very similar to the *Cs* sequence of *D. simulans*. In both the intron's splice site. The predicted peptide length of species we found a long ORF that extends for 1507 bp *Cs* in *D. melanogaster* is 504 amino acids, compared from the intron, determined in *D. melanogaster* by with 544 amino acids in *S. lebanonensis*, if we assume a comparison of our genomic sequence with the cDNA single exon. sequence of Eveleth and Marsh (1987). The longest The regions suggested as protein coding regions are ORF previously proposed is 735 bp. Thus, the coding characterized by somewhat increased codon bias along region of *Cs* is twice as long as previously thought their length (not shown), which is indicative of coding (Eveleth and Marsh, 1987). In addition, the putative regions (Gribskov et al., 1984). Fig. 3 shows the effective amino acid sequence differs from the one previously number of codons, ENC, for six genes, including *Cs* suggested for *D. melanogaster* in several stretches, some and the flanking *amd* and *Ddc* genes, in the three species, as long as 30 amino acids. The *Cs* stop codons of the *S. lebanonensis*, *D. melanogaster*, and *D. simulans*. In all three species are in corresponding positions on our three species, codon-use is less biased for *Cs* than for aligned sequences, although several gaps are necessary any of the other genes, although it is rather similar to in order to obtain the alignment (Fig. 2). The alignment that for *amd* (ENC=61 when all codons are evenly of the encoded peptide sequences obtained by translating used,  $ENC = 20$  when only one codon per amino acid the ORF yields 95% amino acid identity between *D.* is used). *melanogaster* and *D. simulans*, and 78% between *Scaptodrosophila* and those two species.

Although the similarity of the inferred coding regions **4. Discussion** is high, this high similarity does not start from the very beginning of the coding region. We are thus unable to *amd*, *Cs*, and *Ddc* are neighboring genes in *D. melano*use sequence comparisons between *D. melanogaster* and *gaster* (Eveleth and Marsh, 1986). *amd* and *Ddc* are *S. lebanonensis* for elucidating the whole length of the quite similar in nucleotide and amino acid sequences, coding regions. This is not surprising because the coding and are paralogous genes arising from an ancient gene segment of the first exon in *D. melanogaster* is very duplication (Eveleth and Marsh, 1986; Wang et al., short, just three codons, according to Eveleth and Marsh 1996). *Ddc* has been sequenced in a number of organisms (1987). We have used several methods to infer the start (Tatarenkov et al., 1999), but the *amd* and *Cs* sequences of the coding region in *S. lebanonensis*, and have applied have been reported only for *D. melanogaster*. the same methods also to *D. melanogaster*. The programs Comparison of the *Ddc* sequences available in GenBank GENIE and FGENED both predict an intron on the with those of *amd* from a number of species (our *D. melanogaster* sequence as detected by Eveleth and unpublished data) suggests that the duplication of these Marsh (1987) by comparing cDNA with genomic DNA. genes occurred well before the split of Lepidoptera and They also predict the first short exon postulated by Diptera and may predate the divergence of Protostoma Eveleth and Marsh (1987). The first eight nucleotides and Deuterostoma, which occurred more than 600

In order to test these inferences, we sequenced the *Cs* consisted of a continuous single exon in the past, and The corrected sequence of *Cs* in *D. melanogaster* is sequence in *S. lebanonensis* as a result of mutation in



Fig. 3. Codon usage bias in six genes in *S. lebanonensis*, *D. melanogaster*, and *D. simulans*. A larger effective number of codons (ENC) indicates lesser codon usage bias.

MYA, before the Cambrian (Jackson, 1990). If this still not possible to answer when *Cs* arose between *amd* inference is correct, *amd* should be present in many and *Ddc*. Comparisons with species distantly related to animal phyla, unless it has been obliterated, or has *Drosophila* are necessary, such as remote dipterans, evolved beyond recognition. The physical proximity of other insects, and Crustacea. These comparisons will *amd* and *Ddc* most likely traces back to the time of the also help in dating the time of the ancestral duplication original duplication of these genes, but the presence of leading to *amd* and *Ddc*. *Cs* codes for a product of 544 *Cs* between them is enigmatic. It could reflect the amino acids in *S. lebanonensis*, but 504 amino acids in survival by a gene that was contiguous to the duplicated *D. melanogaster* and *D. simulans*. The larger size than gene that led to *Ddc* and *amd*, or it may have been previously proposed of the postulated coding region in inserted there at a much later time. The position in *D. melanogaster* is robust, because in addition to such diverse animal groups of the orthologous genes to these characteristics of coding regions as increased GC bias three might permit us to resolve this issue.  $\qquad \qquad$  and certain codon preferences, the predicted polypep-

ment of genomic DNA comprising partially the flanking *melanogaster* and *D. simulans*; 75% between them and *Ddc* and *amd* genes and an intermediate region in *D. S. lebanonensis*), which would be unexpected in non*melanogaster*, *D. simulans*, and *S. lebanonensis*, a dro- coding regions. sophilid species which diverged from *D. melanogaster* The great proximity of the three genes, *amd*, *Cs* and about 60–65 MYA ( Kwiatowski et al., 1994, 1997). A *Ddc* in *D. melanogaster* is quite unusual (see discussion PCR fragment of similar length was obtained from by Eveleth and Marsh, 1987; but see Okuyama et al., several other Drosophilids. The fact that the region has 1997). The correct coding region of *Cs* that we have remained unchanged in several independent lineages now determined in *D. melanogaster* makes the gene during the last 30–40 Myr may be indication of its density even more spectacular, with the stop codons of functional importance. The comparison of the region *Ddc* and *Cs* genes being only 366 bp apart. Our study between *amd* and *Ddc* in *S. lebanonensis* and *D. melano-* shows that the tight packing also occurs in *S. lebanogaster* has revealed the presence of the *Cs* gene in *S. nensis*, in which the *amd* stop codon is just 686 bp from *lebanonensis*, as it was already known in *D. melanogaster* the *Cs* start codon, and the stop codons of *Cs* and *Ddc* (Eveleth and Marsh, 1987). Moreover, this *Cs* gene is are only 722 bp apart from one another (Fig. 1). The also present in *D. simulans*, where the sequence and suggestion that mutagenic silence of the *Cs* may have exon–intron arrangement is extremely similar to our occurred in *D. melanogaster* as a consequence of evolusequence of *D. melanogaster* (but importantly different tionarily recent modifications in the gene's structure at a few nucleotide sites from a previously published (Eveleth and Marsh, 1987) becomes unconvincing, given sequence; see Eveleth and Marsh, 1987). However, it is that *Cs* has remained tightly packed with *Ddc* and *amd*

We have amplified and sequenced a 5.8 kb-long frag-<br>tides exhibit high sequence similarity (95% between *D*.

Table 1

Number of non-synonymous (n-syn) and synonymous (syn) substitutions per site + SE, and their ratio (n-syn/syn), between *Drosophila melanogaster*, *D. simulans*, and *Scaptodrosophila lebanonensis* at six nuclear genes. The sequences of *Adh* are from Russo et al. (1995); *Gpdh* from Kwiatowski et al. (1997); *Sod* from Kwiatowski et al. (1994)

		melanogaster-simulans	melanogaster-lebanonensis	simulans-lebanonensis
Amd	n-syn	$0.010 + 0.004$	$0.105 + 0.013$	$0.099 + 0.012$
	syn	$0.151 + 0.027$	$1.313 + 0.174$	$1.372 + 0.188$
	ratio	0.066	0.080	0.072
$\mathbb{C}$ s	n-syn	$0.027 + 0.005$	$0.212 + 0.016$	$0.217 + 0.016$
	syn	$0.149 + 0.022$	$1.562 + 0.204$	$1.467 + 0.186$
	ratio	0.181	0.136	0.148
Ddc	n-syn	$0.003 \pm 0.002$	$0.066 + 0.010$	$0.064 \pm 0.010$
	syn	$0.064 + 0.018$	$1.239 + 0.175$	$1.155 + 0.156$
	ratio	0.047	0.053	0.055
Adh	n-syn	$0.002 \pm 0.002$	$0.101 + 0.016$	$0.103 + 0.017$
	syn	$0.052 + 0.021$	$0.802 + 0.122$	$0.765 + 0.117$
	ratio	0.038	0.126	0.135
Gpdh	n-syn	$0.000 + 0.000$	$0.012 + 0.005$	$0.014 + 0.005$
	syn	$0.060 + 0.019$	$1.296 + 0.251$	$1.194 + 0.209$
	ratio	0.000	0.009	0.012
Sod	n-syn	$0.000 + 0.000$	$0.113 + 0.020$	$0.108 + 0.019$
	syn	$0.114 + 0.037$	$1.508 \pm 0.473$	$1.938 \pm 1.030$
	ratio	0.000	0.075	0.056

*lebanonensis* and *D. melanogaster*. The structure and the PCR fragments are of similar length). sequence of this region have remained essentially iden-<br>Eveleth and Marsh (1987) failed to recover *Cs* phenotical in *D. simulans* and *D. melanogaster*, that is for typic mutants in their extensive mutagenesis screens and some 2.5 Myr (we have not investigated the region suggested that this implies that the *Cs* function is not upstream of the *Cs* coding sequence in *D. simulans*, but essential or that *Cs* RNA does not encode a protein, as

for a considerable time, at least 60–65 Myr in *S.* it also seems quite similar with respect to length, since



Fig. 4. Fourier transform of the substitution pattern in four genes: *amd*, *Ddc*, and *Cs* are from *D. melanogaster*, *D. simulans*, and *S. lebanonensis*, *hsr-omega* is from *D. melanogaster* and *D. pseudoobscura*. A dominant substitution frequency of 1/3 is revealed for the exon sequences of *amd*, *Ddc*, and *Cs*, while no predominant peak is observed for the two exons of the non-coding *hsr-omega* (a peak very close to the *y*-axis is due to nonspecific correlations and is largely diminished when gaps are eliminated from the alignment; data not shown).

described for *hsr-omega* in *Drosophila* (Fini et al., 1989). We propose, however, that *Cs* retains protein-encoding capacity. Thus, the ratio of non-synonymous to synonymous substitutions between *D. melanogaster* and *D. simulans*, or between the later two and *S. lebanonensis*, is much less than 1 (Table 1). In a complementary analysis (which may be more appropriate for a comparison between distantly related species, such as *S. lebanonensis* with respect to *D. melanogaster*/*D. simulans*, because of the possible saturation at synonymous sites) the pattern of substitutions also indicates a proteinencoding capacity for *Cs*. If so, *Cs* should have a dominant peak of periodical substitutions every third base. Periodicity in DNA sequencies can be unveiled using Fourier analysis (Tsonis et al., 1991), which we have investigated using an algorithm by Cooley and Tukey (1965). A clear dominant substitution frequency of 1/3 (i.e. every third position) is observed for *amd*, *Ddc*, but also for *Cs*, while it is not for *hsr-omega* (Fig. 4), although the percentage of substitutions is similar in the four genes: 27.8%, 21.6%, 36.6%, and 24.4%, respectively. Nevertheless, there is some indication that the selective constraints may be somewhat lower for *Cs* than for other genes. Thus, the ratio of non-synonymous to synonymous substitutions is higher in *Cs* than in the other five genes (Table 1). Additionally, the number of non-synonymous substitutions per site is higher in *Cs*. Moreover, several gaps, some as long as 10 codons, are needed to align the *Cs* sequences of the three studied species, whereas only one or three gaps are required in *amd* and *Ddc*.

The hypothesis of lesser functional constraints imposed on *Cs* is furthermore supported by analysis of codon usage bias, which is lowest in *Cs* for all three Fig. 5. Amino acid substitutions along *amd*, *Cs*, and *Ddc* between *D.* Drosophilidae that we have studied (Fig. 3). Irrespective *melanogaster*, and *D. simulans* or *S. lebanonensis*. The three boxes of the mechanism underlying the natural selection on<br>silent sites (e.g. rates of protein elongation, translational<br>accuracy), codon usage is typically most biased in highly<br>accuracy), codon usage is typically most biased i expressed genes with high functional constraints (Shields indicate regions where the alignment was not feasible owing to the et al., 1988; Moriyama and Hartl, 1993; Akashi, 1994; absence of at least one sequence. Substitutions are shown by vertical<br>Morivama and Powell 1997). Note however, that lines: short when the amino acid replacements are co Moriyama and Powell, 1997). Note, however, that lines: short when the amino acid replacements are conservative (D/E, although the and power as in  $C_2$  is not as pronounced  $K/R/H$ , N/Q, S/T, I/L/V, F/W/Y, or A/G, according although the codon usage bias in Cs is not as pronounced<br>as in  $\frac{K}{N}$ ,  $\frac{$ not untypical for *Drosophila*. Particularly, ENC in *Cs* is down when they occur in *D. melanogaster*, and up for the compared rather close to that in the neighboring *amd*. Earlier sequence. Numbers indicate the number of amino acid residues. A observations that aciden use the is in C<sub>n</sub> is very weak scale is at the bottom. observations that codon usage bias in *Cs* is very weak compared with other *D. melanogaster* genes (Eveleth and Marsh, 1987; Stathakis et al., 1995) may have arisen reveals statistically significant non-random clustering of from mistakes in the previously published sequence substitutions  $(P<0.01)$  in all three genes in the compariof *Cs*. son between *D. melanogaster* and *S. lebanonensis*, both

tions in the deduced amino acid sequences of *amd*, *Cs*, (short lines in Fig. 5). Interestingly, the non-randomness and *Ddc* (Fig. 5). The three proteins show seemingly is more pronounced for all three proteins when conservadifferent distribution of the substitutions, with *Cs* tive substitutions are not considered (i.e. *P* values are appearing as the most homogeneous. However, the smaller). A graphical plot of the distribution of the unmodified variance test of Goss and Lewontin (1996) segment sizes between substitutions shows an excess,



We have studied the spatial distribution of substitu-<br>including and excluding the conservative substitutions

in *Ddc* and *amd*, and of contiguous substitutions in *Cs* suggesting that the split of *Cs* from the common (data not shown). A remarkable area of low contrained ancestral gene is very ancient, perhaps predating the evolution is the carboxyl end of *amd*, while the central diversification of the major multicellular kingdoms. areas of *amd* and *Ddc* appear to be the most constrained ones. In *Cs* it is difficult to distinguish areas of low interspecific variation, and the non-random distribution of substitutions is probably due to an excess of runs of **5. Conclusions** contiguous substitutions, as mentioned above.

As shown above *Cs* appears to be a protein-encoding (1) Gene order and direction of transcription of the gene. Consequently, we have conducted an extensive *amd*, *Cs*, and *Ddc* genes are the same in *S. lebanonensis* search of GenBank for sequences that would be similar and *D. melanogaster*. *Cs* is very closely packed with the to *Cs*, and have at least six sequences that are distantly neighboring *Ddc* and *amd* genes in *S. lebanon* to *Cs*, and have at least six sequences that are distantly related to *Cs*, although more similar than expected by well as in *Drosophila*. chance. Fig. 6 displays the protein alignments with the (2) The *Cs* gene codes for a longer product than had two most similar sequences: a corticosteroid-binding been previously suggested for *D. melanogaster*. The protein in the yeast *Candida albicans*, and a polyamine length of the deduced protein is 544 amino acids in *S.* oxidase in maize, *Zea mays*. Although the similarity of *lebanonensis* and 504 amino acids in *D. melanogaster*. these sequences to *Cs* is not very high, they are surely In *S. lebanonensis* the protein is encoded by a single homologous. First, the probability of the sequence sim-<br>ORF, while in *D. melanogaster* the coding sequence is ilarity observed is in both cases  $P < 10^{-6}$ . Moreover, the interrupted by a short intron. alignment encompasses large segments of the genes: (3) There is heterogeneity in substitution pattern about 90% of the *Cs* gene in *S. lebanonensis* and 95% between and within *amd*, *Cs*, and *Ddc*. *Ddc* appears to in *D. melanogaster* and virtually the whole extension of be the most constrained gene of the three, especially its the coding regions in the genes of *Candida* and *Zea*. central area. *amd* is less constrained, with a highly Other sequences of about the same length and with variable carboxyl end and a more conserved central similarity nearly as large include amine oxidase in a fish area. *Cs* is affected the most by the substitution process, (P49253); protoporphyrinogen oxidase in tobacco with runs of contiguous substitutions along its whole (Y13466); and proteins of unknown function with sim- length. ilarities to monoamine oxidase and protein kinase in (4) Compared with some other nuclear genes, the *Caenorhabditis* (z78198, locus 1491653) and *Arabidopsis* Drosophilidae *Cs* sequences exhibit higher number of (G2244987). No single sequence that is particularly non-synonymous substitutions, higher ratios of non-

compared with random distribution, of large segments these sequences are approximately equally similar to it,

close to the *Cs* gene could be singled out; instead, all synonymous to synonymous substitutions, and lower

Corticosteroid-binding protein in yeast (Candida albicans)			Polyamine oxidase in maize (Zea mays)		
leban: 55 yeast: 2	SAKONTOIVVIGAGLAGLSAAQHLLRHGFRS---TIVLEATDRYGGRVNS-----KRFGD 106 $S$ ++T++++IGAG++GL AA+ +L $F$ + $+V+EA$ +R GGR+ + $+ G$ SRTKSTKVLIIGAGVSGLKAAETILSKSFLTGDDVLVVEAQNRIGGRLKTTDTSOSKLGI 61	leban: 50 maize: 23	QYNLESAKQNTQIVVIGAGLAGLSAAQHLLRHGFRSTIVLEATDRYGGRVNSKRFGDTYC 109 0+ +A +++V+GAG++G+SAA+ L G ++LEATD GGR++ F OHGSLAATVGPRVIVVGAGMSGISAAKRLSEAGITDLLILEATDHIGGRMHKTNFAGINV 82		
	leban: 107 TYCELGAKWVNMNIDGAHNTIYELLRNAEGLRKOLKO-----RECANYVHTOGREVPPNM 161 $Y + LGA W + D + N + + N + GL$ $++++$ T EVP K	leban: 110	ELGAKWVNMNIDGAHNTIYELLRNAEGLRK------QLKQRECANYVHTQGREVPPNMVE 162 $L_{\odot}$ ELGA WV $G$ N I + + + + LR		
veast: 62	NY-DLGASWFH---DSLNNIVLNHMIN-DGLLDDEKDVYFDDKDLKTFSSTG--EVP--- 111	maize: 83	ELGANWVEGVNGGKMNPIWPIVNSTLKLRNFRSDFDYLAQNVYKEDGGVYDEDYVQKRIE 142		
	leban: 162 VELIDMQFRQLCRGFKVSEKVKSGGDLHVLDNVMAYFKTESEKLVGHSYPDPEKRALARE 221 $++D + +$ $VL+++Y + + +G$ PD R + +		leban: 164 LIDMOFROLCRGFKVSEKVKSGG--DLHVL----------------DNVMAYFKTESEK 204 $D V + Y + K + E$ L D $G$ K+S $+ G$ $D+ +L$		
	veast: 112 -- IVDKKLNR-------------------VLEDIEKYIQLYFNRNLG--VPDLSLRDIVAQ 149		maize: 143 LADSVEEM---GEKLSATLHASGRDDMSILAMQRLNEHQPNGPATPVDMVVDYYKFDYE- 198		
	leban: 222 IFOS----LFKEFSSILGCCLEYVNI------EHITS--CPVOOELRPLYVPTGLDNVLD 269 $F + + +E$ $G + Y +$ $+I+$ $+$ R L $G + + +$		leban: 205 LVGHSYPDPEKRALAREIFOSLFKEFSSILGCCLEYVNIEHITSCPVQQELRPLYVPTGL 264 $P$ $+L$ $+$ $+$ $F$ $+F$ $+$ $+$ $E$ $+$ $+Y$ G		
	veast: 150 YFEKYNRLITEEOREYCGRMMRYLEFWFGISWDRISGKYAVTTHOGRNLLNKKGYGYLVE 209		maize: 199 ----FAEPPRVTSLQNTVPLATFSDFGDDVYFVADQRGYEAV-----------VYYLAGQ 243		
	leban: 270 TLTQHISKEQLQTGKPVGSIQWQTLSDFGAPTSPLPQERKCVACLDGTLYSADHIICTLP 329 $+L + I + L + PV I + DG$ $+R$ $V$ $++G$ $D++I$ T+P		leban: 265 DNVLDTLTQHISKEQLQTGKPVGSIQWQTLSDFGAPTSPLPQERKCVACLDGTLYSADHI 324 D + I +LQ K V I++ G T V D ++YSAD++		
	yeast: 210 SLAKRIPESSLLLEEPVNKII-RNNKDAG--------KRVLVETINGLQIFCDYLIVTVP 260		maize: 244 YLKTDDKSGKIVDPRLOLNKVVREIKYSP----GGVT---------VKTEDNSVYSADYV 290		
	leban: 330 LGVL----KNFSAILFKPALPLEKLOAIRNLGYGNPVKIYLAYKRPISRWLKSNLRPLGA 385 $+L$ $+$ $+I$ $++P LP$ $++I$ $+$ $+G$ $K+$ $+R$ $K + +$		leban: 325 ICTLPLGVLKNFSAILFKPALPLEKLOAIRNLGYGNPVKIYLAYKRPISRWLKSNLRPLG 384 $KT+L + R$ + + LGVL++ I FKP LP K++AI $W + R$		
	yeast: 261 QSILLLEESSPYSIKWEPKLPQRLVESINSIHFGALGKVIFEFDRIFWDNSKDRFOIIAD 320		maize: 291 MVSASLGVLQS-DLIQFKPKLPTWKVRAIYQFDMAVYTKIFLKFPRKF--WPEGKGR--- 344		
	leban: 386 OLGKDEPAITVNGROERLWTOOVVEISOLPSSOHVLEIRVGGGYYDEIEKLPDVTLLEOI 445 $+ + + \nabla + + + + \nabla$ $+ +E$ PD D.		leban: 385 AQLGKDEPAITVNGRQERLWTQQVVEISQLPSSQHVLEIRVGGGYYDEIEKLPDVTLLEQ 444		
	yeast: 321 HTDGDLSRELTELPKPFTYPLFAVNFGRVHNGKASLVILTQAPLTNYLETHPDQAWQYYQ 380		maize: 345 ------EFFLYASSRRGYYGVWOEFE-KOYPDA-NVLLVTVTDEESRRIEQQSDEQTKAE 396		
	leban: 446 TALLROCLRNRLVPYPOALLRSNWSTSACYLGGRPYFSTTSSARDV----ORLAEPLGDI 501 $L + + + + + P$ P + ++W+T+ G $T$ $D+$ $E$ LG $+$		leban: 445 ITALLROCLRNRLVPYPOALLRSNWSTSACYLGGRPYFSTTSSARDVQRLAEPLGDIAPT 504 I +LR+ + VP +L W + Y G + + + +L P+G +		
	yeast: 381 PMLQKLSINDEPIPDPINTIVTDWTTNPYIRGSYSTMYTNDDPSDLIISLSGDFEDLGIL 440		maize: 397 IMQVLRKMFPGKDVPDATDILVPRWWSDRFYKGTFSNWPVGVNRYEYDQLRAPVGRV--- 453		
	leban: 502 APTLLFAGDATALKGFGTIDGARTSGIREAORIID 536 $P$ + FAG+ T +G G + GA SGI A I++		leban: 505 LLFAGDATALKGFGTIDGARTSGIREAORIID 536 $F G+ T+ G + GA SGI A++I+$		
	yeast: 441 EPYIKFAGEHTTSEGTGCVHGAYMSGIYAADCILE 475		maize: 454 -YFTGEHTSEHYNGYVHGAYLSGIDSAEILIN 484		

Fig. 6. Similarity between the deduced *Cs* protein in *S. lebanonensis* and two other proteins: corticosteroid-binding protein in the yeast *Candida albicans* (PIR: A47259), and polyamine oxidase in maize (GenBank: AJ002204). The numbers at the two ends of each row refer to amino acid sites in the proteins. Identical amino acids are shown by letters in the middle rows; crosses indicate functionally similar amino acids.

so highly constrained as the other genes.<br>
(5) The Cs protein exhibits statistically significant<br>
sequence similarity to other proteins, such as some Marsh, J.L., Erfle, M.P., Leeds, C.A., 1986. Molecular localization,<br>
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