

MHC Class II Pseudogene and Genomic Signature of a 32-kb Cosmid in the House Finch (*Carpodacus mexicanus*)

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Large-scale sequencing studies in vertebrates have thus far focused primarily on the genomes of a few model organisms. Birds are of interest to genomics because of their much smaller and highly streamlined genomes compared to mammals. However, large-scale genetic work has been confined almost exclusively to the chicken; we know little about general aspects of genomes in nongame birds. This study examines the organization of a genomic region containing an *Mhc* class II B gene in a representative of another important lineage of the avian tree, the songbirds (Passeriformes). We used a shotgun sequencing approach to determine the sequence of a 32-kb cosmid insert containing a strongly hybridizing *Mhc* fragment from house finches (*Carpodacus mexicanus*). There were a total of three genes found on the cosmid clone, about the gene density expected for the mammalian *Mhc*: a class II *Mhc* β -chain gene (*Came-DABI*), a serine-threonine kinase, and a zinc finger motif. Frameshift mutations in both the second and third exons of *Came-DABI* and the unalignability of the gene after the third exon suggest that it is a nonfunctional pseudogene. In addition, the identifiable introns of *Came-DABI* are more than twice as large as those of chickens. Nucleotide diversity in the peptide-binding region of *Came-DABI* ($\Pi = 0.03$) was much lower than polymorphic chicken and other functional *Mhc* genes but higher than the expected diversity for a neutral locus in birds, perhaps because of hitchhiking on a selected *Mhc* locus close by. The serine-threonine kinase gene is likely functional, whereas the zinc finger motif is likely nonfunctional. A paucity of long simple-sequence repeats and retroelements is consistent with emerging rules of chicken genomics, and a pictorial analysis of the "genomic signature" of this sequence, the first of its kind for birds, bears strong similarity to mammalian signatures, suggesting common higher-order structures in these homeothermic genomes. The house finch sequence is among a very few of its kind from nonmodel vertebrates and provides insight into the evolution of the avian *Mhc* and of avian genomes generally.

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Long DNA sequences provide one source of the genomic information that will revolutionize biology, yet cosmid-scale (25–40 kb) or longer DNA sequences are still almost exclusively confined to model organisms and microbial pathogens. Whereas several nonmodel mammal species are the focus of large-scale mapping and genome projects (O'Brien et al. 1999), cosmid-scale sequences of nonmammalian organisms are available only from chickens, Japanese quail, zebrafish, and pufferfish. We expect the genomic features gleaned from such models to predict aspects of the genomes of related species in their respective clades. Nonetheless, the full diversity of genomic structures will not be appreciated until a much larger number of genomes and DNA sequences from nonmodel species are investigated. To this end we have been investigat-

ing cosmid-scale sequences of birds, with particular attention to the immunologically important major histocompatibility complex (*Mhc*) region (Edwards et al. 1999). Here we report on the first cosmid-scale sequence from a songbird, the house finch (*Carpodacus mexicanus*), a member of the large clade Passeriformes that includes over half of all avian species (Edwards 1998).

The *Mhc* is a multigene family found thus far only in jawed vertebrates. *Mhc* genes have yet to be found in jawless fish or any lineage more ancient (Kandil et al. 1996), although allorecognition genes potentially related to *Mhc* genes have been found in tunicates (Magor et al. 1999). The primary function of the *Mhc* is to present foreign peptides from pathogens to T cells during the adaptive immune response (Klein 1986). *Mhc* genes are the most polymorphic genes found in vertebrates, and much research has been directed toward understanding their evolutionary dynamics, with particular emphasis on possible rela-

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tionships between *Mhc* diversity and parasite resistance (Klein et al. 1993; Parham and Ohta 1996; Edwards and Hedrick 1998). Molecular interactions of *Mhc* genes and pathogen peptides may lead to a “molecular arms race” with recurring bouts of coevolution between the host and the parasite (the Red Queen hypothesis; Van Valen 1973; Hamilton 1982), or *Mhc* diversity may be elevated because of disassortative mating between *Mhc*-dissimilar individuals (Penn and Potts 1998, 1999). This latter view is not inconsistent with a role for *Mhc* genes in defending hosts against parasites. Chickens have provided particularly powerful models for implicating *Mhc* genes in resistance to infectious disease (Briles and McGibbon 1948; Schat et al. 1994; Kaufman and Salamonsen 1997). Structurally, the coding regions of avian *Mhc* genes have many similarities to those of other vertebrates with both class I genes responsible for immune responses to intracellular parasites and class II genes that bind extracellular parasites (Kaufman et al. 1990; Shiina et al. 1999b). The chicken *Mhc* is also known to possess class III *Mhc* genes such as factor B that are involved in the complement system of the cellular immune response (Nonaka et al. 1994). The complete sequence of the chicken *Mhc* (B complex) is an order of magnitude smaller and much more densely packed with genes than mammalian *Mhcs* (Kaufman et al. 1999a,b).

Avian genes and genomes are thought to be subject to a variety of selective pressures imposed by flight. For example, the small size of avian genomes and chicken introns compared with those of mammals and the low frequency of simple-sequence repeats are thought to be due to selection for small cell size to optimize the high metabolic demands for flight (Tiersch and Wachtel 1991; Hughes and Hughes 1995; Primmer et al. 1997). The high gene density of the chicken *Mhc* is thought to reflect similar flight-induced genomic streamlining (Parham 1999). Birds are also known to possess a higher frequency of GC-rich isochores than mammals (Bernardi et al. 1997). However, the global similarities and differences of avian and mammalian genomes are still poorly understood. The concept of a “genomic signature” has emerged in recent years as one way to describe the higher-order structure, mutational biases, and selection pressures underlying genomes as revealed in the frequencies of DNA words of different length observed in long DNA sequences (Karlin and Burge 1995). Novel quantitative and qualitative methods permit description of the genomic signature in ways that are virtually independent of global base composition and isochores structure, thereby providing a common metric by which to compare genomes of different species (Jeffrey 1990, 1992). Deschavanne et al. (1999) reported that, contrary to intuition, the signature of an entire genome or of sev-

eral megabases of a species’ DNA can be accurately captured in just a few dozen kilobases and that a species’ genomic signature is surprisingly robust to the isochores or gene region from which the DNA sequence for the signature is sampled. The few genomic signatures from mammals that have been published reveal, among other things, the characteristic deficiency of CG dinucleotides that had been noted in earlier analyses of mammalian sequences (Deschavanne et al. 1999), and we were curious to see how an avian genomic signature compared with those of mice and humans.

We have been studying the *Mhc* region from house finches (*Carpodacus mexicanus*), both because house finches are well studied ecologically and because they represent an understudied avian lineage with respect to *Mhc*. House finches, a model species for studies of sexual selection and parasite resistance (Hill 1991; Luttrell et al. 1996; Dhondt et al. 1998), are socially monogamous songbirds found throughout the United States (Hill 1991, 1993). House finch *Mhc* class II B genes have been partially characterized via Southern blot analysis and by examining expressed genes through RT-PCR (Edwards et al. 1995a,b, 1999). The house finch *Mhc* contains fewer hybridizing elements when probed with a conspecific cDNA probe than do other songbird species, but we know nothing of house finch *Mhc* genes at the genomic level, nor anything about the noncoding genomic context of *Mhc* genes in any songbird (Edwards et al. 2000; Wester Dahl et al. 1999). To add a phylogenetic perspective to chicken *Mhc* studies, and to characterize the genomic signature of house finches, we sequenced a 32-kb cosmid insert (HFcos10A) from a house finch that strongly hybridized with a house finch class II B clone.

RESULTS

Base Composition and Repeated Elements of Cosmid HFcos10A

The sequence of the cosmid clone HFcos10A was 31,936-bp long (Fig. 1). The GC content averaged 56.9% over the entire cosmid and varied from 33.1% to 70.1% in moving windows of 500 bp. The highest GC content was found in coding regions and the lowest just preceding these regions (Fig. 1). There were a total of 35 exons and 8 genes predicted by Genemark (Fig. 1B); these predictions also tended to occur in regions of relatively high (>60%) GC content. There were 16 simple sequence repeats (microsatellites) found. However, only a single microsatellite was longer than five repeat units (AGA₈). In addition, we identified two LINE elements beginning at positions 802 and 22,221 using the program RepeatMasker (A. Smit and P.

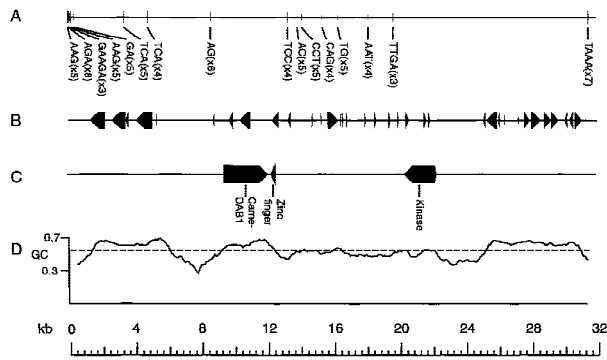


Figure 1 Organization of cosmid HFc0s10A containing simple sequence repeats (A), predicted exons found by the program GeneMark (B), three genes identified using various analysis packages (see text for details) (C), and the GC content plotted with a moving window size of 500 bp and offset length of 50 bp using the program PercentGC (D). Direction of arrows in B indicates the transcriptional orientation of each gene. Dashed line in C is the average GC content over the entire cosmid.

Green, unpubl.), but these LINEs were not verified by subsequent BLAST searches, suggesting that they may not be legitimate. However, their short lengths (32 and 261 bp) are within the range for truncated LINEs found in mammalian *Mhcs* and genomes (Yamazaki et al. 1999).

Structure and Diversity of *Came-DAB1*

SeqHelp identified an *Mhc*-like gene that had the highest GenBank alignment scores with other class II B *Mhc* genes from songbirds; we designate the gene *Came-DAB1* as per *Mhc* nomenclature rules. *Came-DAB1* contains the first three exons expected for typical *Mhc* class II B genes, but the final three exons are not identifiable (Fig. 2). Figure 3 shows an alignment of exons 2 and 3 of *Came-DAB1* with homologous sequences from chicken and red-winged blackbird. There are at least three frameshift mutations located in the second and third exons of *Came-DAB1*. At 439 and 777-bp, respectively, the sizes of introns 1 and 2 are 2.11 and 8.93 times bigger than the corresponding chicken introns. We found only a single *Mhc* gene in >30 kb of cosmid sequence, suggesting a low density of *Mhc*-like sequences in this region of the house finch genome compared to the chicken *Mhc*.

The nucleotide diversity of *Came-DAB1* is consistent with the non-functional nature of the gene. *Mhc* peptide binding regions (PBRs, exon 2 in the case of class II B genes) tend to have a large number of non-synonymous differences as compared with silent changes. We reconstructed the inferred haplotypes for exon 2 from direct sequencing diploid PCR products using the program HAPINFER (Clark 1990; Fig. 4). We used the reconstructed haplotypes to estimate the pattern of nucleotide substitution in the PBR of *Came-*

DAB1 and for phylogenetic analysis of this exon. Figure 4 identifies the different haplotypes that occur in each individual and the specific base found at each segregating site. The program HAPINFER was unable to resolve the phase in two of the individuals, but all subsequent analysis on exon 2 used the inferred haplotypes and not the direct sequence data. The number of substitutions per nonsynonymous site (d_n) is low compared with the number per silent site (d_s) for *Came-DAB1* and compared with typical functional genes from chickens (Fig. 5).

Table 1 describes the overall diversity of *Came-DAB1* using the statistics Π (average pairwise difference) and $\Theta = 4N_e\mu$, where N_e is the effective population size and μ is the neutral mutation rate. Levels of genetic diversity of exon 2 are more similar to the levels for the nonclassical *B-LBIII* gene of chickens than to either of the classical chicken genes (*B-LBI* and *B-LBII*) or a polymorphic blackbird class II B gene (Table 1; Garrigan and Edwards 1999). In particular, we found much lower values of Θ and Π at the *Came-DAB1* locus than found in similar surveys of polymorphic chicken *B-LBI* and *B-LBII* genes. Tajima's D statistic (Tajima 1989), which tests for neutrality in DNA sequence data, was negative for both the finch and the *B-LBIII* locus in chickens, a suggestion of a gene under directional selection although the values are not significantly different from the neutral expectation ($P >> 0.05$). *B-LBI*, *B-LBII*, and the blackbird gene (*Agph-DAB1*) all had positive values for Tajima's D , consistent with balancing selection. HAPINFER was unable to resolve inferred haplotypes from exon 3 data from *Came-DAB1*, perhaps because of a number of alternate homozygotes found in the direct sequences. Nonetheless, we can still examine diversity (Θ) using the number of segregating (polymorphic) sites (Waterson 1975). As expected from a pseudogene, the values of Θ for exon 3 are similar to those of exon 2 (Table 1). Moreover, these values are somewhat higher than the neutral Θ found in other vertebrates such as humans. For example, Grimsley et al. (1998) found Θ values of 0.0262 for *HLA-H*, an *Mhc* pseudogene linked

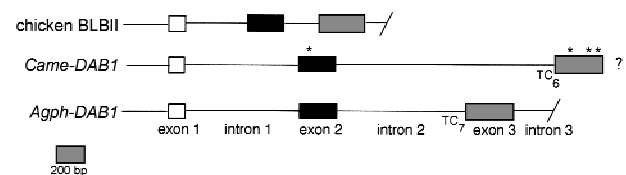


Figure 2 *Mhc* class II B structure in house finches (*Came-DAB1*), red-winged blackbirds (*Agph-DAB1*; Edwards et al. 1998), and chickens (*B-LBII*; Zoorob et al. 1990). The asterisks represent frameshift mutations indicating the nonfunctional nature of *Came-DAB1*. The shaded exon represents the polymorphic second exon encoding the peptide binding region. Intron and exon sizes are to scale. Dashes signify that the gene continues downstream.

A, exon 2

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1          11          21          31          41          51          61          71          81          91          100
Came-DAB1  GTGTTCCAGG GGATGAGATT GAA-GAGTGT CACTTGTGA  ACGGCACGGA GAAGGTGAGG TTCGTTGAGA GGTTCATCTA CAACCCGGAG CAGTCCCTGA
HFcDNA    A.....CT.AA .TCC.....CACC.....C.....G.....
Chicken   T.C...TCTT ACGGTGTGA. ATTT.....C .A..CC.....C..CG.....ATC.....CAA.....C.....TC.C
|
101        111        121        131        141        151        161        171        181        191        200
Came-DAB1  TATTTGGAGC GGACGTGGGG GGTACATGGG GGTTCATCGC CTATGCGGGG ATGAATGCCA AGCGCTCTAA CAGGCACCGG GTTCACATGG AGTACTACCG
HFcDNA    G..C.....G.....CAC.....C..C.....TAT.....CTG.....CG.....ACATAC.....A.A...
Chicken   AC..C.....G.....AAA.....CCGAT.CAC. GCTG.AT.. TATC.G.TAG .AAT..GG...CG..G.C .AGTTTC...A..CGAAT
|
201        211        221        231        241        251        261
Came-DAB1  GGCTTTGGTG GACACGCCGT GCTGGCGAAA CTAAGAGCTT TATGCCCCGT TCACAACGCT GGAGCCGC
HFcDNA    A..GC.....G.T.C..C..AC.....G.G..CCCG.....TC....A.CGC...A
Chicken   AA.GAA.....G.T.C..C..AC.....G.GA. GTC.AGT.C. ...GTT.CA .AG.A..

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B, exon 3

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1          11          21          31          41          51          61          71          81          91          100
Came-DAB1  TCCCCCC-AG TGTGTGCTCT TCTGGGTTGC CC--TCGAC CTCGACCACC GGGCCACAGC GCGTGCCTGT CTCGGTCAAT GATT--TCT ACCCGCCCCA
HFcDNA    C.....C.....A.....C.....C.....CCC.....G.....G.....T.....G.....C.....T.....C.....C
Chicken   .GAG..C.A G.....G.....C..GC.C..A. AG.....G.....T.....T.....AAA..GA..T..GCG..A...G.C. GGCGCT...GC.GG.
|
101        111        121        131        141        151        161        171        181        191        200
Came-DAB1  CATCCAGGGG AGTGTGTTCC AGGGCCAGCA GGAGCTCTCA GACCACATGG TGGCCACAGA CATGGTCCCC GAGAACAGGG ACTGGAC-TA CCAGTCCCTG
HFcDNA    G.....T.....G.....G.....G.....G.....G.....C.....G.....T.....G.....C.....G.....G.....
Chicken   G..G...T.....A.....T.AA.GE..G .GAG.A.G .G.G.E.G....T.....G.....G..A.G.AG .T.....G.....G.....G.....G.....
|
201        211        221        231        241
Came-DAB1  CTGCTGG AAA-CCACC CCAGGCGAGG CTCACCTACA CCAGCCAG
HFcDNA    .G..CGG..G.C..G..G.....T.....
Chicken   .G.C.GT...G.G.C..G..GA..G...G TGT...G.

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Figure 3 Alignment of *Came-DAB1* exon 2 (A) and exon 3 (B) to a highly polymorphic chicken gene (*B-LB1*; Zoorob et al. 1993) and a house finch cDNA sequence (Edwards et al. 1995a). Deletions are indicated in the sequences by a – mark.

to the highly polymorphic *HLA-A* gene and reported that these Θ values were an order of magnitude higher than the background Θ for humans. A neutral Θ for house finches is not known and neutral intron diversity in some seabirds, which are known to have large population sizes, are about the same (H. Walsh, pers. comm.). Although it is not definitive we think that this value of Θ would be high for finches at a neutral locus.

Origin of *Came-DAB1*

Came-DAB1 exon 2 and 3 sequences were easily aligned to those of other functional and nonfunctional *Mhc* genes. Phylogenetic trees of *Came-DAB1* using the neighbor joining method (Saitou and Nei 1987), the inferred haplotypes for exon 2, and direct sequence data for exon 3 in general exhibit a strong trend toward clustering of sequences by species (Fig. 6a,b). The phylogenetic reconstructions of both exon 2 and exon 3 place the sequences from HFc10A closest to the house finch sequences obtained for the polymorphism study. However, both the exon 2 and exon 3 trees suggest that the sequences from the *Came-DAB1* locus are not the closest relatives of the expressed cDNA sequences of house finch class II B genes from Edwards et al. (1995a). For exon 3 the *Came-DAB1* sequences are most closely related to sequences from the Bengalese finch (*Lonchura striata*, *Lost*; family Estrildidae; Vincek et al. 1995). The branch lengths leading to the Bengalese finch in the exon 2 tree are deep compared to the branches leading to

the *Came-DAB1* sequences, and the bootstrap value supporting monophyly of finch and blackbird sequences is not high. However, the Bengalese finch and the *Came-DAB1* sequences cluster strongly (100%) for exon 3 despite the fact that these two “finches” are in different taxonomic families.

Non-*Mhc* Genes

A serine-threonine kinase gene detected by SeqHelp –8340 bp from *Came-DAB1* is predicted to be transcribed in the opposite direction as *Came-DAB1*. The sequence was aligned to two homologous sequences using a BLAST search. The highest similarity genes were members of the Ste20/PAK family from *Xenopus*, which is involved in the arrest of oocytes at G_s /prophase of the first meiotic

cell cycle and prevention of apoptosis (Faure et al. 1997) and a *Drosophila* homolog of the serine-threonine kinase PAK gene that has a potential function in focal adhesion and colocalizes with dynamic actin structures (Harden et al. 1996). The house finch gene has three alignable exons, with the first and third exons slightly shorter than the genes mentioned above (not shown). Both a start and stop codon are found within a few amino acids of these features in the other sequences.

A zinc finger motif is found just downstream of *Came-DAB1* and also runs in the opposite transcriptional orientation. The total length of the motif is 79 amino acids, whereas the two sequences examined from GenBank with which it had the highest similarity were 574 and 1207 amino acids long. There was only a single alignable exon containing two C2H2 motifs [C2H2 motifs are tandemly repeated domains commonly found in zinc finger proteins and comprise one of the most common gene families in the human genome (Becker et al. 1995)], apparent from the BLAST search. A BLAST search of the 100 nucleotides upstream of this region did not yield any convincing hits.

Genomic Signature

We investigated the genomic signature implied by the 32-kb house finch sequence using the pictorial chaos-game representation (CGR) algorithm of Descavanne et al. (1999), in which the frequencies of different DNA words are depicted by varying shades, from black (most

Individual Number (inferred haplotypes)	3	20	22	38	58	61	77	100	107	129	134	153	154	155	165	208	215	225	227	234	248	250	251	254	256	
1(*08,*09)	C/G	T	A	A/T	G	C/T	C/G	A/G	A	G	A/G	A/C	C/T	C/G	G	C	G	G	A/G	A/G	G	G	T	C/T	A	C
2(*01,*02)	G	T	A	A/G	G	C/T	C	A	A	G	A	A	C/T	G	C	G	G	C	A	A/G	G	T	C	A	G	G
3(*01,*01)	G	T	A	A	G	C	C	A	A	G	A	A	T	G	C	G	C	G	C	A	G	G	T	C	A	G
4(*01,*03)	G	A/T	A/T	A	G	C/T	C/T	A	A	G	A	A	T	G	C	G	G	C	A	A	G	G	T	C	A	G
5(*06,*07)	C/G	T	A	A/T	C/G	C	C	C/G	C/G	A/G	A	A/C	C/T	C/G	C/G	G	G	C	A	A	G	C/G	T	C	A/G	G
6(unresolved)	G	T	A	A	G	C	C	A	A	G	A	A	T	G	C	G	C	A	A/G	C/G	T	C	A	A	G	
10(*01,*04)	G	T	A	A	G	C	C	A	A	G	A	A	T	G	C	A/G	G	C	A	A/G	C/G	C/T	C	A/C	G	
11(*01,*01)	G	T	A	A	G	C	C	A	A	G	A	A	T	G	C	G	G	C	A	A	G	G	T	C	A	G
13(*01,*05)	C/G	T	A	A/T	G	A/C	C	A	A	G	A	A	T	G	C	G	C/G	C	A	A/G	G	T	C	A	G	
16(unresolved)	G	A/T	A/T	A	G	C	C	A	A	G	A	A	T	G	C	G	G	C	A	A	G	G	T	C	A	G
<i>Came-DAB1*01</i>	G	T	A	A	G	C	C	A	A	G	A	A	T	G	C	G	G	C	A	A	G	G	T	C	A	G
<i>Came-DAB1*02</i>	G	T	A	A	G	T	C	A	A	G	A	A	C	G	C	G	G	C	A	A	G	T	C	A	G	
<i>Came-DAB1*03</i>	G	A	A	A	G	T	T	A	A	G	A	A	T	G	C	G	G	C	A	A	G	T	C	A	G	
<i>Came-DAB1*04</i>	G	T	A	A	G	C	C	A	A	G	A	A	T	G	C	A	G	C	A	A	G	C	C	C	G	
<i>Came-DAB1*05</i>	C	T	T	T	G	A	C	A	A	G	A	A	T	G	C	G	C	C	A	A	G	T	C	A	G	
<i>Came-DAB1*06</i>	G	T	A	A	G	C	C	C	C	G	A	A	T	G	C	G	G	C	A	A	G	G	T	C	A	C
<i>Came-DAB1*07</i>	C	T	A	T	C	C	C	G	G	A	A	C	C	G	G	G	C	A	A	G	C	T	C	G	G	
<i>Came-DAB1*08</i>	G	T	A	A	G	C	C	A	A	G	A	A	T	G	C	G	G	C	A	A	G	G	T	C	A	C
<i>Came-DAB1*09</i>	C	T	A	T	G	T	G	G	A	G	G	C	C	C	C	G	G	G	G	G	G	G	T	T	A	C

Figure 4 List of variable sites as inferred from HAPINFER (Clark 1990) for exon 2 sequences. (Top) Unresolved sequences with reconstructed genotypes in parentheses; (bottom) the reconstructed alleles. HAPINFER was unable to resolve phase two of the individuals (see text for details). Column numbers refer to the site where the polymorphisms occur and are consistent with the alignment from Fig. 3.

frequent) to white (least frequent). We investigated the frequency of DNA words of two (dinucleotides), five, and eight letters (Fig. 7) on both strands of the finch sequence. In the CGR method, a square image is divided into four quadrants signifying the four nucleotides. The pixel signifying the frequency of all DNA 'words' of any length ending in a given nucleotide occurs in that nucleotide's quadrant. Each quadrant is in turn divided into four quadrants that signify the nucleotide occurring in the second to last position of words. These secondary quadrants occur in the same positions relative to one another as do the original quadrants, and so on, until the appropriate number of pixels (4^n , where n is the number of letters in the words being investigated) is achieved. In this way certain large-scale features of the sequence examined can emerge. For example, dark diagonals indicate stretches composed solely of purines or pyrimidines, and empty patches in upper left quadrant of the upper right quadrant indicate a deficiency of words containing CpG dinucleotides.

The genomic signature of the house finch exhibits strong signals on the diagonals for five- and eight-letter words, indicating a high frequency of purine and pyrimidine stretches (Fig. 7). It also exhibits a notable CpG depletion for all three word lengths, as indicated by the pale regions in the upper left quadrants of all three upper right quadrants, as well as a deficiency of TA dinucleotides, as indicated by the pale lower right quadrant of all three lower left quadrant (Fig. 7). This latter result occurs despite the presence of several TA-rich microsatellites (albeit short ones; Fig. 1). We conducted a quantitative analysis of the five-letter word frequencies. We found that two words are never met in the sequence or on its complementary strand—TACGC and GCGTA, both of which contain the two counter-

selected dinucleotides CG and TA). Consistent with the G+C rich nature of sequence, the most frequent five-letter words are CCCTG, CAGGG, GGGGA, TCCCC, GGGGG, CCCCC, GGCCA, TGGCC, CCCAG, CTGGG, CCCCT, AGGGG, CCCCCA, and TGGGG, all of which occurred between 255 and 292 times, about 5–6 times the median of the distribution of five-letter words in the entire sequence.

DISCUSSION

Status of Sequenced Genes on Cosmid HFc0510A

In conjunction with new sequences from red-winged blackbirds (Edwards et al. 2000), the cosmid we have sequenced is the first cosmid-scale sequence determined for any avian species other than chicken. It thus provides a glimpse into the genomic architecture of the most species-rich clade of birds, Passeriformes, as well as insight specifically into the structure of regions containing *Mhc* genes. At about one identified gene per 10 kb, the gene density of the region we have sequenced is more similar to that of the mammalian *Mhc* than to the chicken B complex (MHC Sequencing Consortium 1999; Kaufman et al. 1999b). Only one of the 35 exons predicted by Genemark corresponded accurately to the manually identified exons, the zinc finger, and neither *Came-DAB1* nor the serine-threonine kinase genes were predicted accurately. We therefore suspect that many of these predictions are spurious. Although we cannot be sure that the house finch sequence occurs in the same genomic region as polymorphic and presumably functional *Mhc* genes, that is, in the canonical house finch *Mhc* (Edwards et al. 1995a, 1999), *Mhc*-containing regions of this gene density have not yet been reported for any avian species and our estimate of avian gene density is only the second (after the chicken B complex) for any bird based on cosmid sequences. *Came-DAB1* is among a very few avian *Mhc* genes sequenced at the genomic level (Guillemot et al. 1988; Zoorob et al. 1990; Edwards et al. 1998; Kaufman et al. 1999b). It has none of the attributes of a classical *Mhc* gene. The gene does not have a high rate of nonsynonymous substitutions, an expected signature of a functional *Mhc* gene under balancing selection, nor high levels of total diversity (Π and Θ). All indications are that *Came-DAB1* is a pseudogene. There are frameshift mutations in two of the three identifiable exons (including the PBR-encoding

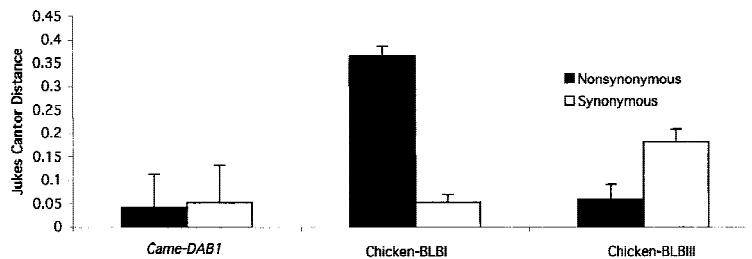


Figure 5 Comparison of the number of nonsynonymous (d_n) and silent (d_s) substitutions per site within house finches for exon 2 in *Came-DAB1* ($n=9$), a functional chicken gene ($n=12$), and a “nonclassical” class II chicken gene ($n=3$). These comparisons were made with a number of chicken sequences downloaded from GenBank (Zoorob et al. 1993) and the nine inferred haplotypes from house finches obtained through PCR amplification, direct sequencing, and analysis using HAPINFER as described in text. The d_n and d_s values were calculated by the Jukes-Cantor method of Nei and Gojobori (1986) using the MEGA software package (Kumar et al. 1993).

exon) and the sequence similarity to other *Mhc* genes declines after the third exon. We used the method of Miyata and Yasunaga (1981) to estimate the time since loss of function for this pseudogene. This method uses the difference in the d_n/d_s ratio in comparisons of the focal pseudogene, a functional homolog (ingroups) and an outgroup sequence to estimate time since loss of function in the pseudogene. We used the sequences from exon 2 of a functional blackbird gene Edwards et al. (1998) as the outgroup and *Came-DAB1* and cDNA sequences from the house finch (Edwards et al. 1995a) as ingroups in this analysis. The estimated time since loss of function of *Came-DAB1* is $0.9 T_0$, where T_0 is the time since divergence of the blackbird and house finch. DNA-DNA hybridization studies (Sibley and Ahlquist 1990) place this split at about 50 million years ago (MYA), making the time since loss of function of *Came-DAB1* at ~45 MYA. This ancient date is consistent with the d_n/d_s ratios at *Came-DAB1*, which appear to have reached base substitutional equilibrium. The

ratios were not significantly different than one, a pattern expected in a pseudogene after a long period of neutral evolution.

The two other genes found on the cosmid have not been found near *Mhc* genes of other birds (Kaufman et al. 1999a,b; Shiina et al. 1999b). However, genes found in the same broadly defined multigene families have been found inside the *Mhcs* of chickens and other vertebrates. For instance, the *RING3* gene is a kinase that is found in the class II region of mammals, chickens, and frogs (Kaufman et al. 1999a). However, the house finch kinase is clearly not similar to *RING3*. Genes similar to the house finch serine-threonine kinase gene are involved in protein phosphorylation (Kruse et al. 1997) and the zinc-finger protein is a widespread transcription factor motif in eukaryotic genomes (Struhl 1989). There is no reason to expect that these genes are involved in the antigen presenting process.

Multigene Family Evolution

Because *Came-DAB1* was easily aligned with other avian and vertebrate *Mhc* genes, and because phylogenetic analysis clearly showed that *Came-DAB1* clustered with other expressed songbird *Mhc* genes, *Came-DAB1* can justifiably be called an *Mhc* gene. In designating *Came-DAB1* as an *Mhc* gene, we differ with Kaufman et al. (1999a), who suggest that only *Mhc*-like genes that reside in regions homologous to the chicken B-complex and are expressed and functionally important should be designated as such (Miller et al. 1994). We prefer a genealogical definition of *Mhc* genes, rather than a functional one. The fact that *Came-DAB1* does not cluster specifically with functional chicken *Mhc* genes (Fig. 6a,b) does not bear on the question of

Table 1. Comparison of Nucleotide Diversity of *Came-DAB1*, Three Chicken Genes, and a Red-Winged Blackbird Gene *Agph-DAB1*^b

Gene	<i>Came-DAB1</i>		Chicken exon 2 ^a			<i>Agph-DAB1</i> ^b
	exon 2	exon 3	<i>B-LBI</i>	<i>B-LBII</i>	<i>B-LBIII</i>	exon 2
No. of sequences examined	9	10	12	11	3	17
Total no. of sites	260	287	270	270	270	89
No. of segregating sites(s)	25	32	138	54	41	22
Θ	9.19	11.48	44.47	18.43	22.36	6.32
Θ per site	0.04	0.04	0.17	0.07	0.08	0.07
II	8.89	—	47.99	22.64	22.33	8.90
Nucleotide diversity (II per site)	0.03	—	0.18	0.08	0.08	0.10
Tajima's <i>D</i>	-0.17	—	0.36	1.08 ^c	-0.01	1.99 ^c

Data from both exons 2 and 3 from *Came-DAB1* are shown. Θ is calculated using the finite sites method of Tajima (1996). All relevant statistics are discussed in Methods.

^aFrom Zoorob et al. (1993).

^bFrom Garrigan and Edwards (1999).

^c $P < 0.03$.

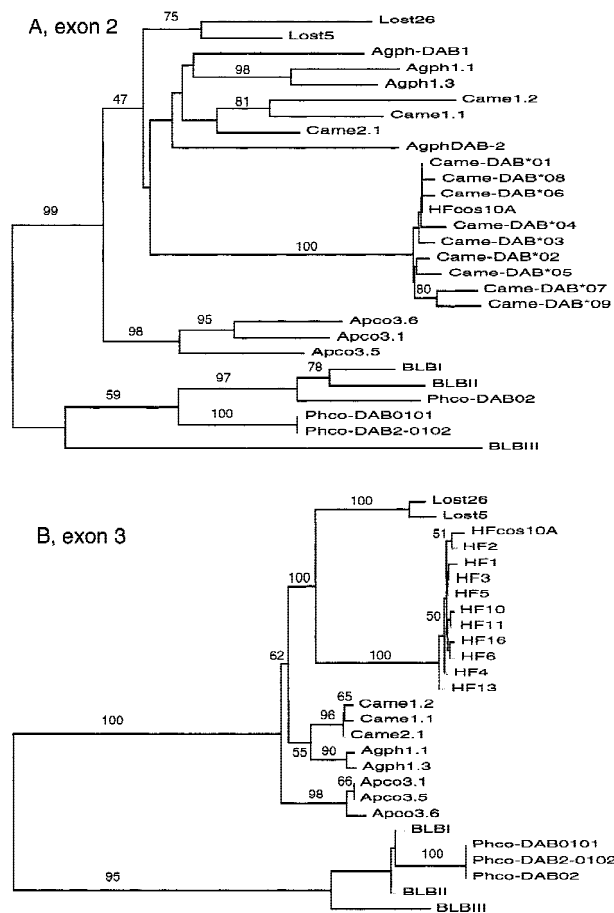


Figure 6 Phylogenetic analysis of exon 2 (A) and exon 3 (B) sequences from different avian species (Lost=Bengalese finch, Vincek et al. 1995; Apco=scrub jay, Came=house finch, Agph=red-winged blackbird, Edwards et al. 1995a; HFcos10A=house finch cosmid 10A, HF=unresolved exon 3 sequences; Phco=ring-necked pheasant, Witzell et al. 1999; BLB=chicken, Zoorob et al. 1993). The trees shown are neighbor-joining trees (Saitou and Nei 1987) using a Tamura-Nei (1993) distance. The numbers above the branches are bootstrap scores with 1000 replicates. The trees are rooted with the pheasant and chicken sequences used collectively as an outgroup.

whether it resides in the genomic region homologous to the chicken B-complex, given the possibility of functional *Mhc*-genes being dispersed in birds, as occurs in zebrafish (Bingulac-Popovic et al. 1997). Given what we know of multigene family evolution in avian *Mhc* genes (see below), we expect most songbird *Mhc* genes to form clusters separately from those of chickens, regardless of whether they are expressed or not.

Given that we can align *Came-DAB1* to other *Mhc* sequences and perform a phylogenetic analysis, we are justified in discussing multigene family evolution in the avian *Mhc*, as we have in the past (Edwards et al. 1995a; 1999). That *Came-DAB1* is a pseudogene supports the idea of a birth and death model of *Mhc* evolution (Ota and Nei 1994; Nei et al. 1997). The birth and death model predicts that there is frequent gene

duplication and pseudogene formation in multigene families. Our phylogenetic analysis addressed some of the hypotheses about the mode of *Mhc* multigene family evolution in birds—whether a concerted evolution (Witzell et al. 1999) or a divergent evolution model is most prevalent. The data for both exons 2 and 3 largely support a concerted evolution model because the predominant pattern is clustering by species. However, exon 3 sequences from *Came-DAB1* are more closely related to genes from other species (Bengalese finch, family Estrilididae) than they are to the presumably functional house finch genes obtained from cDNA. Either *Came-DAB1* is orthologous to the genes from the Bengalese finch (exon 3) or the absence of stabilizing selection acting on *Came-DAB1* has masked its phylogenetic signal through homoplasy (exon 2). Homoplasy is expected to be a problem more for the second exon of *Mhc* genes because of the increased likelihood of base substitutional saturation caused by balancing selection and, more importantly, the scrambling effects of recombination and gene conversion. Exon 3 is a better indicator of gene history because it is not under the diversifying selection pressures, and the strong clustering of the Bengalese finch with *Came-DAB1* supports its orthology with the *Lost* sequences.

Genomic Signature

The HFcos10A sequence contained several microsatellites, mostly with five or fewer repeat units (Fig. 1). The low density of long microsatellites, with only two repeats with a total length >20 bp in 30 kb of sequence, is consistent with a low density of simple sequence repeats found in another survey of avian microsatellites (Primmer et al. 1997). These researchers found an average of one microsatellite of >20 bp total length per 39 kb, a low density compared to human DNA (1 microsatellite per 6 kb; Beckmann and Weber 1992). For the human *Mhc* class I region the number of microsatellites is ~1 per every 2 kb (Shiina et al. 1999a).

The CGR genomic signature exhibited by the house finch sequence, the first reported for birds, displays a number of similarities to mammalian signatures and raises a number of predictions for signatures of other avian genomes. The house finch signature bears a number of similarities to other homeothermic vertebrates thus far examined (e.g., mouse and human). The deficiency of CpG dinucleotides, as well as the relatively high frequency of purine or pyrimidine runs (diagonals) suggests that these may be features of vertebrate genomes generally. The deficiency of CpG is intriguing given evidence for a much higher density of genes and CpG islands in the chicken than in the mammalian genome (McQueen et al. 1996) and the higher frequency of GC-rich isochores in birds compared with mammals (Bernardi et al. 1997). The deficiency of TA dinucleotides and the longer words em-

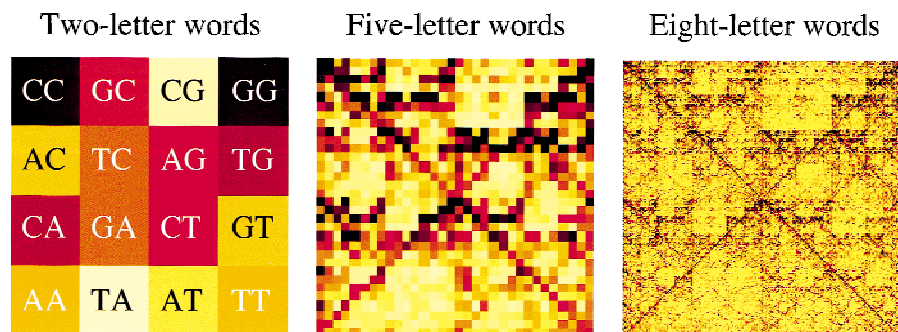


Figure 7 Genomic signature of house finch DNA for two-, five-, and eight-letter words based on the 32-kb cosmid sequence. Dark pixels indicate high-frequency words and light pixels indicate low-frequency words. The 16 (4^2) words in the leftmost panel are indicated for convenience. The five-letter signature contains 4^5 words (pixels) and the eight-letter signature 4^8 . The fact that the orientation of the four nucleotides is the same at all scales in the image, i.e., within any given quadrant of any size, contributes to the fractal nature of the image.

bedding this and CG words is a novel feature that is not as pronounced in the mammalian signatures that have been investigated to date by the CGR method (Deschavanne et al. 1999). We note that the region we have sequenced is somewhat more GC-rich (and TA-depauperate) than mammalian sequences, as expected for birds (Bernardi et al. 1997). However, this TA deficiency of the genomic signature remains even after the effects of global base composition have been removed (data not shown). Some of these features could be the result of particular directional mutation pressures resulting from the high metabolic rates and high body temperatures of birds; such features are known to influence the mutational spectrum and base composition of animal mitochondrial DNA (Martin 1995). Pettigrew (1994) suggested that flying vertebrates should have elevated levels of A and T nucleotides because of higher metabolic demands. However, analysis by Van Den Bussche et al. (1998) showed that flying mammals such as bats do not have higher AT levels than other mammals. Our results suggest that birds also may not show the elevated AT levels predicted by Pettigrew (1994). Although the sequence we have analyzed is only 32 kb, we suspect it will capture many of the features of avian genomic signatures based on longer sequences, in part because of the patent similarities of the signature to those of non-*Mhc* regions in mammals; the signature for 8-mers may be less precise than those for shorter words because the frequency of each of the 4^8 8-mers may not be captured accurately even in 32kb. The house finch signature therefore can be tested for generality by analyzing sequences from avian species that are less well studied in this regard, such as the chicken.

METHODS

Samples

Cosmid 10A was isolated from a cosmid library as per Edwards

et al. (2000). All birds used in the polymorphism screen are the same as those used in Edwards et al. (2000) as well as four more birds from the same Alabama population. All birds are unrelated. All sequences have been deposited in GenBank with accession numbers AF205032 and AF241546–AF241565.

Sequence Assembly

We sequenced the clone containing the most strongly hybridizing band as revealed by a Southern blot analysis probed with a partial (exon 1–4) RT-PCR product from a house finch class II B *Mhc* gene (see Edwards et al. 1998, 1999b for details on cosmid library construction and

screening). We then sonicated the cosmid clone, subcloned the sonicated fragments into an M13 vector, and prepared multiple clones using Qiagen Prep Kits in a 96-well format. We sequenced at random 830 subclones on an ABI 373 cycle sequencer with Dye-terminator chemistry using a modified M13-T7 (5'-TGCTGCAGGTCGACTCTAG) vector primer. These sequences were aligned and assembled into contigs using the program PhredPhrap (Ewing and Green 1998; Ewing et al. 1998). We visualized the assembled contigs using the program Consed (Gordon et al. 1998), designed primers for the end of each contig, and connected the contigs by walking. This method of primer design was also used to improve regions of low sequence quality after the first round of sequencing.

The final contig was analyzed for sequence similarity with GenBank sequences using the program SeqHelp (Lee et al. 1998). We also predicted open reading frames and exons using the program GeneMark (Lukashin and Borodovsky 1998). GeneMark uses a Markov model to statistically search for splice signals and start codons generated from a matrix derived from empirical observations. In our case a chicken matrix was used: simple sequence repeats using Sputnik (C. Abajian, unpubl.) and more complex repeats using RepeatMasker (A. Smit and P. Green, unpubl.). The genomic signature was analyzed for words of 1–8 letters using the methods outlined in Deschavanne et al. (1999). The frequencies of all words of varying lengths was determined after concatenating both strands into one sequence. This action was taken because the genomic signature is strand dependent (see Deschavanne et al. 1999 for details).

Polymorphism Analysis

We designed locus-specific PCR primers to amplify the sequences from exons 2 and 3 of *Came-DAB1* (exon 2: 5' HF10AEX2F-GCTGTGTCCTGCACTCACA, 3' HF10AINT2R.1-GCAGGGTCCGAGGGGAC; exon 3: 5' HF10AINT2F.1-CTGATTCCAGTGTGTCCCCA, 3' HF10AINT3R.1-CCAGTGGCTCTCCCAGTG). This was accomplished by comparing the cosmid *Mhc* sequence to previously published cDNA sequences (Edwards et al. 1995a) and maximizing the areas of discrepancy. We directly sequenced 10 individuals (both strands) for all of exons 2 and 3 using the forward and reverse PCR primers as well as two internal sequencing primers both in the 5' and 3'

directions (exon 2: 5' HF10EX2F.2-GAGAGGTTTCATCTCAACCG, 3' HF10EX2R.2-AGCTCGTAGTTTCGCCAGC; exon 3: 5' HF10AEX3F.2-CTCTCTCCTCCCTCACAG, 3' HF10AEX3R.2-CCGGGGCTCCCCATAT). These sequences were then aligned and a consensus sequence for each individual was generated using the alignment program Sequencher (Gene Codes). Sequences were checked manually and examined for the presence of heterozygous sites. We then used the program HAPINFER (Clark 1990) to generate haplotypes. These haplotypes were then used to generate estimates of the d_n/d_s ratios (the number of nonsynonymous to synonymous changes) using the Jukes-Cantor method (Nei and Gojobori 1986) and to infer Tajima's D (Tajima 1989), the number of segregating sites (s), and Θ (Watterson 1975) using a program DNAPOLY written by Dan Garrigan (unpubl.).

Phylogenetic Analysis

We included the 9 inferred haplotypes for exon 2 and the 10 individuals yielded from direct sequencing for exon 3 in our analysis. We were unable to infer the haplotypes for exon 3 and therefore used diploid sequences in our phylogenetic analysis—technically a violation of standard phylogenetic analysis, but one with likely little effect given the low level of variability. Also included in these analyses were sequences downloaded from GenBank from the Bengalese finch (*Lonchura striata*, Vincek et al. 1995), chicken (*Gallus gallus*, Pharr et al. 1998), scrub jay, and red-winged blackbird (Edwards et al. 1995a). These sequences were aligned using the program Sequencher to lengths of 260 bp (exon 2) and 287 bp (exon 3). We used the neighbor-joining method (Saitou and Nei 1987) and a Tamura-Nei (1993) distance method for all phylogenetic analysis. The resulting trees were rooted using the chicken and pheasant sequences as outgroups. A total of 1000 bootstrap replicates (Felsenstein 1985) were completed to determine the robustness of phylogenetic groupings.

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