Vertebrate genomes are mosaics of isochores—namely, of long (>300 kb), compositionally homogeneous DNA segments that can be subdivided into a small number of families characterized by different GC levels. In the human genome (which is representative of a number of mammalian genomes, and, more broadly, of the genomes of warm-blooded vertebrates), the compositional range of isochores is 30%-60% GC, and five families of isochores have been identified: two GC-poor families, L1 and L2, together representing 62% of the genome, and three GC-rich families, H1, H2, and H3, representing 22%, 9%, and 3%, respectively (the remaining 4% of the genome is formed by satellite and ribosomal DNA). Gene concentration is strikingly nonuniform, being highest in the H3 isochrome family, lowest in the L1+L2 families, and intermediate in the H1+H2 families. The H3 family corresponds to T(elomeric) bands of metaphase chromosomes, and the L1+L2 families correspond to G(iemsa) bands, whereas R(everse) bands comprise both GC-poor and GC-rich isochores. The compositional distributions of large genome fragments, of exons (and their codon positions), and of introns are correlated with each other. They represent compositional patterns and are very different between the genomes of cold- and warm-blooded vertebrates, mainly in that the former are much less heterogeneous in base composition and never reach the highest GC levels attained by the latter. Only relatively small compositional differences are found among the genomes of either cold- or warm-blooded vertebrates. Compositional patterns allow one to define two modes in genome evolution: a conservative mode, with no compositional change, and a transitional (or shifting) mode, with compositional changes. The conservative mode can be observed among either cold- or warm-blooded vertebrates. The transitional mode comprises both major and minor compositional changes. In vertebrate genomes, the major changes are associated with the appearance of GC-rich and very GC-rich isochores in mammalian and avian genomes. Mutational biases play a role in both modes of compositional evolution. According to one viewpoint, the fixation of compositionally biased mutations is responsible for the transitional mode of evolution of bacterial genomes; in the conservative mode of evolution of vertebrates, they accomplish their role in conjunction with differences either in chromatin structures that modulate replication errors or in chromatin transcriptional activities that may lead to various extents of repair-DNA synthesis. According to another viewpoint, defended here, selection controls, at the isochrome level, the fixation of compositionally biased mutations, both in the conservative and in the transitional mode of evolution.

Introduction

Vertebrate genomes are mosaics of isochores—namely, of long, compositionally homogeneous DNA segments that can be subdivided into a small number of families.
characterized by different GC levels. Figure 1 displays a scheme of the isochores forming the human genome, which is representative of a number of mammalian genomes and, more broadly, of the genomes of warm-blooded vertebrates. The compositional heterogeneity of high-molecular-weight bovine DNA was discovered 20 years ago (Filipski et al. 1973) by using CsSO₄ preparative density gradient centrifugation in the presence of a sequence-specific DNA ligand, Ag⁺. This heterogeneity concerned the so-called main band DNA and was different from the heterogeneity previously detected in analytical CsCl gradient (Sueoka 1959), which was essentially due to the eight GC-rich satellite DNAs that form 23% of the bovine genome (Filipski et al. 1973; Cortadas et al. 1977; Kopecka et al. 1978; Macaya et al. 1978). The basic properties of isochores (as they were called by Cuny et al. 1981) from the vertebrate genomes were defined by Thiery et al. (1976) and Macaya et al. (1976) and were later studied in further detail (for a review, see Bernardi 1989).

Isochores from the mouse genome were estimated to be >300 kb in size (Macaya et al. 1976). Recent work has shown that, in the human genome, isochore size ranges between 0.36 and >0.7 Mb in the dystrophin gene (Bettecken et al., accepted) and is >1 Mb in the cystic fibrosis locus (Krane et al. 1991). These sizes [also investigated by Ikemura and Aota (1988) and by Ikemura et al. (1990)] are intermediate between those of genes or gene clusters and those of chromosomal bands; they cover the most interesting range for physical and compositional mapping.

In the human genome, the compositional spectrum of isochores is 30%-60% GC, and five families of isochores have been identified—two GC-poor families, L1 and L2, together representing 62% of the genome, and three GC-rich families, H1, H2, and H3, representing 22%, 9%, and 3%, respectively. The remaining 4% of the genome

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**HUMAN GENOME**

**ISOCHORES (>300 Kb)**

![Diagram of human genome isochores](image)

DNA fragments (ca. 100 Kb)

![Diagram of DNA fragments](image)

**GC range 30 - 60%**

Fig. 1.—Scheme of the isochore organization of the human genome. The genome is a mosaic of large (>300 kb) DNA segments, the isochores, that are compositionally homogeneous and belong to a small number of families, GC poor (L1 and L2), GC rich (H1 and H2), and very GC rich (H3). Physical and enzymatic degradation occurring during DNA preparation generates large DNA fragments, currently ~100 kb in size. (Modified from Bernardi et al. 1985)
Consists of satellite and ribosomal DNAs, which can also be visualized as isochores, because of their homogeneous base composition (Bernardi 1989)."
The compositional correlation that links GC levels of third codon positions of human genes with the GC levels of the extended sequences in which the genes are located (fig. 4A) can be used to assess gene distribution in the different isochrome families and to quantify the finding (Bernardi et al. 1985) that gene distribution in the human genome is strikingly nonuniform. This approach (fig. 5) has shown that, while 34% of all genes currently present in gene banks are contained in isochrome families L1 and L2, 38% are contained in H1 and H2, and 28% in H3 (Mouchiroud et al. 1991). If the gene sample used is representative of all human genes, and if account is taken of
FIG. 4.—A, GC levels of third codon positions from human genes, plotted against the GC levels of DNA fractions (blackened circles) or extended sequences (unblackened circles) in which the genes are located. The correlation coefficient and the slope are indicated. The dash-and-point line is the diagonal line (slope = 1). GC levels of third codon positions should fall on this line if they are identical to GC levels of surrounding DNA. The broken lines indicate a ±5% GC range around the slope. (From Mouchiroud et al. 1991) B, Plot of GC levels of third codon positions against GC levels of first + second positions of prokaryotic and eukaryotic genomes. All values are averaged per genome (or genome compartment, in the case of compositionally compartmentalized genomes). (From D’Onofrio and Bernardi 1992)

the different relative amounts of isochore families (see Introduction), gene concentration in H3 would be 16 times higher than that in L1+L2 and 8 times higher than that in H1+H2. These ratios are, however, probably underestimated, because housekeeping genes, which are likely to be more abundant in H3 than in other isochore families (see next section), are currently underrepresented in gene banks (Mouchiroud et al. 1991).

The results just described (1) indicate that increasing gene concentrations are accompanied by increasing GC levels in the genome of warm-blooded vertebrates (the evolutionary process underlying this phenomenon will be discussed later) and (2) independently confirm the classification of isochore families, which was originally based on purely compositional grounds; for instance, gene concentration is low and constant over isochore families L1 and L2 and is highest in isochore family H3. It is interesting that the gradient of gene concentration is paralleled by a series of changes in a number of properties that have functional significance. This will be illustrated by describing the extreme situation found in the GC-richest isochore family, H3. The situation found in the GC-poor isochore families L1 and L2 corresponds to the opposite extreme situation, while that of isochore families H1 and H2 is intermediate.

The Human Genome Core

The isochore family H3 corresponds to a genome compartment endowed with remarkable properties. This family has not only the highest GC level and the highest gene concentrations but also the highest concentrations of CpG doublets (Bernardi 1985), the only potential sites of methylation in vertebrates, and the highest concentrations of CpG islands (Aissani and Bernardi 1991a, 1991b), which are very GC-rich sequences characterized by abundant, unmethylated CpG doublets (Bird 1986).
Since CpG islands (which are located in the 5' flanking sequences of genes; Aissani and Bernardi 1991a, 1991b) are preferentially associated with housekeeping genes (Gardiner-Garden and Frommer 1987), the latter should be more abundant in H3 than in the other isochore families.

The coding sequences of H3 isochore are higher in GC level than is their genomic environment, compared with the coding sequences from other isochore families, especially those from GC-poor isochore (Aissani et al. 1991). Moreover, these genes and their associated CpG islands are characterized by a particular chromatin structure, with nucleosome-free regions, absence or scarcity of histone H1, and acetylation of histones H3 and H4 (Tazi and Bird 1990; also see Aissani and Bernardi 1991a, 1991b). These properties make these chromosomal regions more "open," as also is indicated...
by the sensitivity to nuclease attack of DNA located in Reverse bands (Kerem et al. 1984; also see next section).

The H3 isochore family presumably has the highest level of transcription because of its very high concentration of genes—especially housekeeping genes. It also has the highest recombination rate, possibly because of its open chromatin structure and because of the abundance of repetitive sequences, such as Alu sequences and minisatellites. The very high recombination rate of H3 isochores may also be largely responsible for the much higher rate of karyotypic rearrangements (and speciation) shown by mammals compared with cold-blooded vertebrates (Bernardi, accepted). Indications exist that the H3 isochores may be the main integration regions for the majority of (GC-rich) retroviral sequences (see Rynditch et al. 1991; Zoubak et al. 1992; also see Two Modes of Genome Evolution: The Conservative Mode section, below).

The H3 isochore family has an extremely biased codon usage, a number of codons being absent or very scarce because of the very high GC levels in third codon positions, and an extreme amino acid utilization, which favors amino acids corresponding to codons having only G and/or C in the first two codon positions (D’Onofrio et al. 1991)—namely, arginine (quartet codons), alanine, glycine, and proline—rather than amino acids corresponding to codons with only A and/or T in those positions, such as lysine, or those corresponding to codons having both G/C and A/T in first and second codon positions, such as serine. Finally, the sequences of the H3 family are located in T-bands of metaphase chromosomes (see next section).

The main reason for proposing the name of genome core for the GC-richest isochore family of the human genome is, however, that both the strikingly nonuniform gene distribution described here for the human genome and, in particular, the existence of isochores with very high gene concentrations appear to be shared by all warm-blooded vertebrates and, very probably, by all vertebrates (author’s unpublished data). If the latter point is confirmed, the compositional pattern of warm-blooded vertebrates characterized by GC-rich and very GC-rich isochores must have been superimposed on a preexisting gene concentration pattern that was already present in cold-blooded vertebrates but not characterized by any large differences in GC levels. Another genome feature predating the formation of GC-rich isochores is the early and late replication timing of DNA, which was already present in cold-blooded vertebrates (Almeida-Toledo et al. 1988; Giles et al. 1988; Yonenaga-Yassuda et al. 1988).

Isochores and Chromosomal Bands

A number of findings indicate that GC-poor isochores are located in G(iemsa) bands, whereas GC-rich isochores are located in R(verse) bands of human metaphase chromosomes (for a review, see Bernardi 1989). The correlations between compositional heterogeneity of isochores and chromosomal bands in warm-blooded vertebrates also explain why metaphase chromosomes from cold-blooded vertebrates, whose genomes are characterized by a low degree of compositional heterogeneity, show very poor or no banding (Cuny et al. 1981; Medrano et al. 1988; Schmid and Guttenbach 1988).

At least in the case of R-bands of human chromosomes, the correspondence cannot, however, be a direct one, for the simple reason that GC-rich and GC-poor isochores are in a 1:2 ratio, whereas R- and G-bands are in a 1:1 ratio. An approach developed to solve the problem of the correlations between isochores and chromosomal bands is compositional mapping (Bernardi 1989). This consists in hybridizing probes corresponding to landmarks that are localized, on either a physical map or a chro-
mosome band map, to compositional DNA fractions. This approach allows one to assess the GC level of ~200 kb around the landmarks, if the fractionated DNA is ~100 kb in size.

Compositional mapping, as applied to the long arm of human chromosome 21 (Gardiner et al. 1990), showed that practically all probes for loci present in G-bands hybridized to GC-poor isochores, whereas probes located in R-bands hybridized to either GC-poor or GC-rich isochores. In other words, G-bands are GC poor and at least largely homogeneous in base composition, whereas R-bands are compositionally heterogeneous. The GC-richest region of the long arm of the human chromosome 21 was shown to correspond to the telomeric band, which is a T-band (Dutrillaux 1973)—namely, one of the 20 or so R-bands most resistant to heat denaturation. This observation prompted an analysis of the localization of single-copy sequences from the isochore family H3. As suggested by previous work on the location of genes in T-bands (De Sario et al. 1991; Ikemura and Wada 1991), this analysis showed that these sequences are located at T-bands (Saccone et al. 1992). This result, a first step toward a compositional map of the human karyotype, shows that a DNA fraction, isolated on the basis of its nucleotide composition, corresponds to a well-defined cytogenetic compartment of metaphase chromosomes.

Very recent compositional mapping results on the Xq26-Xqter region (Pilia et al., submitted) confirmed the GC poorness and compositional homogeneity of G-bands, as well as the heterogeneity of R-bands, and revealed isochores of the H3 family even in chromosomal regions, such as the telomeric Xq28 band, which did not correspond to T-bands. It is therefore possible that a number of “thin” T-bands are still escaping cytogenetic detection. If such is the situation, and if one considers that evidence is available to suggest that “intercalary” (nontelomeric) T-bands are the result of telomeric fusions in evolution (Dutrillaux 1979), then the high concentration of genes in telomeres and “ancestral” telomeres becomes a conspicuous property of the vertebrate genome, the implications of which certainly deserve further investigations. Indeed, human telomeres are tightly associated with the nuclear matrix (De Lange 1992), via their TTAGGG repeats forming the terminal 2–30 kb of chromosomes, and with the nuclear envelope (Henderson and Larson 1991).

Two Modes of Genome Evolution
The Conservative Mode

Compositional patterns allow one to define two modes of genome evolution: the conservative mode and the transitional mode. The conservative mode is characterized by the absence of compositional changes, as observed by comparing either large (~100 kb) DNA fragments or homologous exons (and their codon positions) and introns from different genomes. In the case of the mouse/rat comparison, the compositional distribution of DNA fragments is practically identical for the two species, and differences in third codon positions GC from homologous genes are <1% (Bernardi et al. 1988). Only slightly larger compositional differences were found when third codon positions from human and bovine genes were compared (fig. 6). In the latter case, compositional conservation still holds, in spite of the fact that third codon positions cover a 30%–90% GC range, exhibit a nucleotide divergence averaging 20% (without multiple-hit correction), and are from genomes separated for 60–70 Myr.

The simplest explanation for the conservative mode of evolution would be that, for all isochores, GC-to-AT changes are compensated by an equal number of AT-to-
Third codon position

\[ y = 0.975x + 3.075, r^2 = 0.955 \]

Fig. 6.—GC levels of third codon positions of all available pairs of homologous bovine (ordinate) and human (abscissa) genes, plotted against each other.

GC changes. This would raise no problem for isochores with compositions close to 50% GC. The compositional conservation of isochores (and of the corresponding coding sequences) having extreme GC levels is, however, incompatible with a “random” process of mutation and fixation, which would drive GC levels toward 50%. Maintaining those extreme levels requires a fixation of mutations that are biased in opposite directions for GC-poor and GC-rich isochores. This has been explained by proposing that the bias of the replication machinery is modulated by local chromatin states or by chromatin transcriptional activities that may lead to different extents of repair-DNA synthesis (Sueoka 1988), a point discussed later in more detail.

An alternative explanation is that the conservative mode of genome evolution is due to a negative selection against compositional deviations from a narrow GC range (fig. 7A). Negative selection obviously cannot operate at the single-nucleotide level but can do so at a regional (isochore) level (Bernardi et al. 1988). Some compositional divergence appears to be tolerated, but the upper and lower thresholds seem to be quite close. Indeed, even at size levels as small as those of genes, compositional divergence remains low in third codon positions. Perhaps, therefore, cooperative structural changes (which might somehow be compared to phase transitions) take place
in isochores beyond the thresholds, and perhaps they have deleterious functional consequences (on transcription, e.g.), leading to decreased fitness and to negative selection. According to this hypothesis, at least some of the genes present in an isochore that drifted away from its optimal GC level would produce proteins deficient in quantity and/or quality. This hypothesis does not require negative selection to do more than it admittedly does with regard to classical deleterious mutations in genes.

Note that expressed retroviral sequences integrated within the mammalian genome are located in isochores of matching GC level, whereas nonexpressed sequences from the same viruses are located within isochores characterized by nonmatching GC levels (Bernardi 1989; Rynditch et al. 1991; Zoubak et al. 1992). This may suggest an effect of the chromosomal environment on the expression of integrated viral genomes. Proving this, however, requires detailed comparisons of integrated viral genomes that are not yet available.

The Transitional (or Shifting) Mode

The transitional (or shifting) mode of genome evolution is characterized by compositional changes. Major compositional shifts occurred between the genomes of cold- and warm-blooded vertebrates, and minor ones occurred between genomes within each one of these broad classes. Compositional transitions may be observed at the level of DNA fragments, of exons, and of introns but are best studied by compositional comparisons of different codon positions of homologous genes.

If codon positions of homologous genes from cold- and warm-blooded vertebrates
are compared (Bernardi and Bernardi 1991), GC levels of the coding sequences from warm-blooded vertebrates are either equal to or (with very few exceptions) higher than those of their cold-blooded counterparts (fig. 8), so providing direct evidence for directional base changes (Perrin and Bernardi 1987; Bernardi et al. 1988). Such differences are, as expected, much larger in the third position than in the first+second codon positions. The major compositional transitions leading to mammals and birds, although similar, are not identical, the latter attaining slightly higher GC levels in both DNA fragments and third codon positions than does the former (Thiery et al. 1976; Bernardi et al. 1988).

In terms of base composition, the genome of warm-blooded vertebrates appears, in fact, to comprise a paleogenome, characterized by GC-poor isochores that have not changed in composition relative to the corresponding isochores of cold-blooded vertebrates, and a neogenome, characterized by isochores that have become GC rich (Bernardi 1989; see fig. 9). The GC increase in the genome of warm-blooded vertebrates affects only approximately one-third of the genome, which contains, however, at least two-thirds of the genes. Indeed, as already mentioned, GC increases parallel gene concentration (see fig. 5).

Minor compositional transitions occurred among either warm- or cold-blooded vertebrates. In the first case, transitions separate some mammalian orders and families (Thiery et al. 1976; Bernardi et al. 1988; author’s unpublished data). A special case is that of murids, cricetids, and spalacids, which mainly differ from other rodents (as well as from most other mammals investigated) in showing narrower distributions of both DNA fragments and third codon positions (Salinas et al. 1986; Mouchiroud et al. 1987, 1988). Compared with homologous human genes, third codon positions of GC-rich rat and mouse genes are less GC rich, whereas those of GC-poor genes are less GC poor, the order of GC levels remaining largely the same in the two species (Mouchiroud et al. 1987, 1988). Among cold-blooded vertebrates, a number of compositional transitions have been observed (Bernardi and Bernardi 1990b).

![Fig. 8.—GC levels of third codon positions of pairs of homologous genes from human (ordinate) and cold-blooded vertebrates (abscissa), plotted against each other. (From Bernardi and Bernardi 1991; for further details, see the text.)](image-url)
Compositional Genome Transitions

The Mutational Bias Hypothesis

Two different explanations have been provided for the compositional transitions. The first explanation, originally proposed to account for the different composition of prokaryotic genomes, is that compositional patterns (or genome compositions) shift because of directional mutations due to biases in replication/repair enzymes (Freese 1962; Sueoka 1962, 1988, 1992).

This explanation is generally considered to have been demonstrated by the existence of an Escherichia coli mutator strain, mutT (Cox and Yanofsky 1967), which has a mutation rate 1,000-fold greater than the spontaneous mutation rate, which only induces A-to-C transversions and which has been reported to cause an increase in GC by 0.3% after 1,200–1,600 generations. The evidence for such a minute change should, however, be viewed with caution, because the buoyant density difference reported is within experimental error. Moreover, directional mutations found in mutator strains appear to be introduced at a very limited number of hot spots (Yanofsky et al. 1966; Nghiem et al. 1988; Wu et al. 1990) and are, therefore, unlikely to cause overall compositional genome changes. In any case, for this explanation to be satisfactory, the evidence should be provided that genomes with changed composition are at least not at a disadvantage relative to the unchanged genomes. (This evidence is unlikely ever to be obtained, because of the overwhelming effect of the very high mutation rate).

The absence of any direct evidence does not, however, rule out, per se, the mutational bias hypothesis. Other problems exist, however. Some of them are of a general nature; others are specifically associated with compositional transitions in vertebrates.

Along the first line, the mutational bias hypothesis implies that compositional changes are irrelevant, or neutral, as far as genome organization, function, and evolution are concerned and that they can therefore be left to the vagaries of mutations in the replication/repair systems. This cannot be easily accepted if one considers the
correlations between base composition and DNA structure, the functional importance of the latter, and that compositional changes in the genome are accompanied both by changes in codon usage (reaching even the extreme situation of codon substitutions; Osawa et al. 1987) and by changes in amino acid composition in the encoded proteins. For these reasons, it is difficult to accept the interpretation of differences of base composition in bacterial genomes as essentially due to mutational biases as originally proposed (Freese 1962; Sueoka 1962).

Another general argument against the mutational bias hypothesis is that the spread of GC levels of genomes from different species decreases from bacteria to protists, to invertebrates, to cold-blooded vertebrates, and to warm-blooded vertebrates (Bernardi and Bernardi 1990b), indicating that the base composition of the genome is certainly not freely drifting in all living organisms. In fact, base composition rather appears to be generally related to the variety of the intra- and extracellular environments of the organisms under consideration. In vertebrates, the stronger the homeostasis, the narrower the GC spectrum exhibited by genomes from different species.

Additional serious problems exist for the mutational bias hypothesis in vertebrates. Indeed, as shown in figure 10, mutations in the replication/repair machineries, leading to mutational biases from AT to GC, should have happened only twice, in the two reptilian lineages leading to mammals and birds, respectively (or in the ancestral warm-blooded vertebrates), but never in any other of the numerous families of cold-blooded vertebrates, since the latter never gave rise to genomes compositionally patterned like those of warm-blooded vertebrates. Furthermore, the two series of mutational events led to compositional changes only in the isochores that were to form the neogenomes of mammals and birds and not in the more abundant isochores that later formed the paleogenomes of warm-blooded vertebrates. Finally, once attained, the GC levels of different GC-rich isochore families were maintained by mutational biases that were different from the original ones. This scenario, which involves processes concerning thousands of physically separated isochores within each genome, is most unlikely to have occurred essentially through mutational bias. Indeed, the proposal was put forward that different chromatin structures in different regions of the genome might account for the different directions and effects of mutational biases and that different chromatin transcriptional activities may lead to various extents of repair-DNA synthesis (Sueoka 1988). However, this does not explain why the original GC increases only occurred in the genomes of the ancestors of mammals and birds. The basic problems just mentioned also make it unlikely that changes in nucleotide-precursor pools could account for the formation of GC-rich isochores in warm-blooded vertebrates (Wolfe et al. 1989), a hypothesis afflicted by a number of other problems (see Bernardi et al. 1988; Eyre-Walker 1992).

The minor compositional transitions associated with some cold- or warm-blooded vertebrates also argue against the mutational bias hypothesis. In fish genomes, for example, such transitions are unrelated to evolutionary time since the appearance of the order, family, or genus, nor are they related to the number of species within a given order (Bernardi and Bernardi 1990b). Indeed, if compositional changes were only caused by mutations in the replication/repair machineries, compositional divergence would be expected to be larger for older (or larger) groups and smaller for more recent (or smaller) groups, whereas this is not the case. Moreover, the compositional transitions exhibited by fish genomes show extremely different rates, including very high ones (Bernardi and Bernardi 1990b). This casts doubts about the possibility of constructing correct phylogenetic trees if the genes under consideration (and the
FIG. 10.—Scheme of the formation of GC-rich isochore families in the genome of warm-blooded vertebrates, according to the mutational bias hypothesis. Thick arrows indicate the times at which mutations in the replication/repair machineries led to GC increases in certain compartments of the genomes of therapsids and dinosaurs (giving rise to GC-rich isochore families H1, H2, H3, and H4), but not to increases in other compartments (giving rise to isochore families L). Thin arrows indicate further changes in mutational biases, which led to the maintenance of GC-rich isochore families. All changes in mutational biases should be visualized as operating at thousands of physically separated isochores. At the same time, only minor compositional changes took place in the genomes of all other cold-blooded vertebrates (also see text).
isochores harboring them) underwent compositional transitions (Saccone et al. 1990; Bernardi et al. 1992). Finally, if compositional transitions are as frequent as they appear to be on the basis of recent work on mammals (author's unpublished data), the molecular clock (Zuckerkandl and Pauling 1962) may only apply to genes that have not been subjected to compositional transitions.

The Selection Hypothesis

The second explanation for the compositional genome transitions (such as the major transitions that occurred in the genomes of vertebrates) is that directional mutations are fixed through positive selection operating at the isochore level. Under this explanation, mutational biases only provide the mechanism for compositional changes, but selection controls the compositional levels of isochores.

In general, selective advantages associated with compositional patterns of genomes may be elusive, because the patterns are due to many different factors whose interplay is impossible to sort out. For instance, it is not now possible to provide any explanation for the minor transitions that occurred within either cold- or warm-blooded vertebrates. (However, the narrow compositional patterns exhibited by some rodents, such as rat, mouse, and hamster, might correspond to a partial release of the compositional constraints operating on the isochores exhibiting extreme GC levels.)

This problem may, however, be solved in some situations. One may hope to identify a selective advantage if the advantage is dominant and if it can be evaluated against a relatively similar genomic background, a condition fulfilled by vertebrates. In this case, the major split in genome patterns did not occur at a major step in organismic evolution, like the transitions from anamniotes to amniotes or from fishes to tetrapods, or in a gradual way during the evolution from fishes, to amphibians, to reptiles, and to warm-blooded vertebrates but occurred only and precisely at the transition from cold- to warm-blooded vertebrates. This suggests that the major factor that played a role in the change of compositional patterns of the genome might be related to body temperature. This suggestion not only fills the empty space left by the old mutational bias hypothesis and by its modern "chromatin" version, but it can also be tested.

The increase in GC in the genomes of warm-blooded vertebrates makes sense, as far as selective advantages are concerned, because it leads to thermodynamically more stable DNA, RNA, and proteins (see Bernardi and Bernardi 1986 and references quoted therein). Indeed, GC richness increases the thermal stability of DNA not only in dilute solution but also in chromosomes, as shown by R- and T-bandings, two techniques that show that GC-rich and very GC-rich DNAs are increasingly more stable against thermal denaturation than are GC-poor DNAs from G-bands. GC richness also increases the thermal stability of RNAs, because of the increased secondary structures that make transcripts more stable. Finally, it increases the thermal stability of encoded proteins, because it leads to increased levels of amino acids that confer thermal stability (such as arginine, alanine, and glycine) and to decreased levels of amino acids that reduce such stability (such as lysine and serine).

The objection that some thermophilic bacteria have AT-rich genomes has no relevance for the body-temperature hypothesis, in that comparisons of thermophylic and mesophylic bacterial genomes are not warranted when the species under consideration are separated by enormous phylogenetic distances (e.g., when Eubacteria and Archebacteria are compared) and when they exhibit extremely large differences in cell physiology. The same warning applies to organelle genomes, in which other selective
advantages may be predominant. In addition, thermal stabilization of genomes might be due not to GC increase but to DNA methylation or to protein-DNA interactions.

An independent argument favoring the suggestion just presented is the similarity of compositional patterns of mammals and birds, two vertebrate classes characterized by very different genome sizes and that appeared at different geologic times (>200 and ~150 Mya, respectively) and originated from different ancestral reptiles (therapsids and dinosaurs, respectively). So also do (a) the strong compositional heterogeneity of isochores in plants originating from arid climates (such as wheat and maize), which withstand high maximal temperatures, and (b) the weak compositional heterogeneity of isochores in plants from temperate climates (Salinas et al. 1988; Matassi et al. 1989; Montero et al. 1990). In their compositional patterns, the former resemble warm-blooded vertebrates, and the latter resemble cold-blooded vertebrates. Under the mutational bias hypothesis, all these similarities must be regarded as sheer coincidences.

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LITERATURE CITED


with respect to chromosome banding patterns and chromosome numbers, relation between nucleotide sequence data and cytogenetic data. Nucleic Acids Res. 13:1915–1922.


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