Temporal Patterns of Fruit Fly (Drosophila) Evolution Revealed by Mutation Clocks

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Drosophila melanogaster has been a canonical model organism to study genetics, development, behavior, physiology, evolution, and population genetics for nearly a century. Despite this emphasis and the completion of its nuclear genome sequence, the timing of major speciation events leading to the origin of this fruit fly remain elusive because of the paucity of extensive fossil records and biogeographic data. Use of molecular clocks as an alternative has been fraught with non-clock-like accumulation of nucleotide and amino-acid substitutions. Here we present a novel methodology in which genomic mutation distances are used to overcome these limitations and to make use of all available gene sequence data for constructing a fruit fly molecular time scale. Our analysis of 2977 pairwise sequence comparisons from 176 nuclear genes reveals a long-term fruit fly mutation clock ticking at a rate of 11.1 mutations per kilobase pair per Myr. Genomic mutation clock–based timings of the landmark speciation events leading to the evolution of D. melanogaster show that it shared most recent common ancestry 5.4 MYA with D. simulans, 12.6 MYA with D. erecta+D. ersona, 12.8 MYA with D. yakuba+D. teissieri, 35.6 MYA with the takahashii subgroup, 41.3 MYA with the montium subgroup, 44.2 MYA with the ananassae subgroup, 54.9 MYA with the obscura group, 62.2 MYA with the willistoni group, and 62.9 MYA with the subgroup Drosophila. These and other estimates are compatible with those known from limited biogeographic and fossil records. The inferred temporal pattern of fruit fly evolution shows correspondence with the cooling patterns of paleoclimate changes and habitat fragmentation in the Cenozoic.

Introduction

Inference of species divergence times in the fruit fly phylogeny that led to the evolution of Drosophila melanogaster has been hindered by the absence of extensive, reliable fossil records and biogeographic data. Molecular clocks are routinely used as an alternative tool to infer temporal speciation patterns in such cases (Eastal and Oakeshott 1985; Powell 1997; Hedges and Kumar 2003). In fruit fly evolutionary studies, however, it has been difficult to establish reliable molecular clock calibrations and to obtain unbiased time estimation because of the unequal substitution rates among species, even for synonymous substitutions (Eanes et al. 1996; Rodriguez-Trelles, Tarrio, and Ayala 1999; Tatarenkov et al. 1999; Rodriguez-Trelles, Tarrio, and Ayala 2000). Highly expressed genes show extreme preference for certain codons, which often leads to lower estimates of the number of synonymous substitution per site because of the effects of natural selection and estimation biases (Shields et al. 1988; Sharp and Li 1989; Dunn, Bielawski, and Yang 2001).

Furthermore, codon usage biases are known to vary significantly among lineages (Rodriguez-Trelles et al. 1999), which will lead to synonymous substitution rate variation among lineages. This poses severe problems while inferring molecular time scales of fruit fly evolution. For instance, the Hawaiian Drosophila species, for which the divergence time is considered to be the best point for calibrating molecular clocks, show much lower codon usage biases than D. melanogaster (Rodriguez-Trelles et al. 2000); the D. melanogaster Adh gene sequence shows a 36% higher codon adaptation index (CAI; Sharp and Li 1987) than the Hawaiian D. picticornis Adh gene sequence. This is also true even when we compute codon usage statistics independent of the knowledge of the optimal codons, such as the effective number of codons (Wright 1990); D. picticornis has a much higher effective number of codons (47.1) than D. melanogaster (36.1). This difference in codon usage biases is also reflected in the rejection of the homogeneity of the substitution patterns in the third codon positions at a 1% level when the disparity index test (Kumar and Gadagkar 2001) is used.

Because evolutionary divergences now used for building fruit fly molecular clocks directly employ the actual amount of synonymous or nonsynonymous change that has been permitted in sequence evolution, the intrinsic non-clock-like behavior of substitution accumulation has repeatedly impeded those efforts (Thomas and Hunt 1993; Russo, Takezaki, and Nei 1995; Rodriguez-Trelles, Tarrio, and Ayala 2001a, b). Therefore, to build molecular time scales, we need to estimate distances that are independent of the effects of codon usage bias and selection. Here we present a method for estimating mutation distance based on analysis of multiple genes (we refer to this as genomic mutation distance) and use it for inferring timing of major fruit fly speciation events.

Methods

Sequence Data Acquisition

cDNA and protein sequences for D. melanogaster were obtained from http://www.fruitfly.org/sequence/dMfasta.shtml. All available sequences belonging to the genus Drosophila were downloaded from the National Center for Biotechnology Information (NCBI; GenBank). Redundant sequences were identified based on the annotated gene name and the sequence similarity, and were excluded. A BLASTP (Altschul et al. 1997) search was conducted using each D. melanogaster protein...
sequence as a query and the database of all other fruit fly sequences as the target set. For each set of BLAST-hits produced, we constructed an orthologous sequence data set by taking the best hit from each distinct species available. Sequences in each putative orthologous set were then checked for gene name to ensure the use of the same gene; GenBank protein identification numbers (protein id) were matched in case of gene name ambiguities. When multiple sequences were available for the same gene from a species, the longest sequence was chosen. The data for known genes from Bergman et al. (2002) were then added to this collection. Each protein sequence pair (obtained by translating the cDNA sequence) was aligned using Clustal W (Thompson, Higgins, and Gibson 1994), and the corresponding cDNA alignments were generated with the protein sequence alignments as guides. All analyses were conducted by using these pairs of aligned sequences.

Distance Estimation

### Synonymous Distances

We used the number of nucleotide substitutions per fourfold-degenerate site as the measure of synonymous distance to avoid estimation biases from approximations needed to separate synonymous and nonsynonymous sites (Dunn, Bielawski, and Yang 2001; Kumar and Subramanian 2002). A third codon position was considered fourfold degenerate only if it was fourfold degenerate in both sequences compared. To reduce estimation errors, only sequence pairs with 50 or more fourfold-degenerate sites were included. This produced 6,085 sequence pairs from 176 genes. Of these only 2,977 pairs involved in major divergence events that occurred in the lineage leading to D. melanogaster were used. The Tamura-Nei (Tamura and Nei 1993) method was used to correct for multiple hits in order to account for transition/transversion rate and base-composition biases. The Disparity index test (Kumar and Gadagkar 2001) revealed significant base composition differences among lineages as the stationarity of substitution pattern was rejected in 35% pairwise comparisons at a 5% level, confirming significant differences in codon usage among lineages. Therefore, we used the modified Tamura-Nei method (Tamura and Kumar 2002) to account for substitution pattern heterogeneity in fourfold-degenerate sites among lineages.

### Genomic Mutation Distances

For computing genomic mutation distances, we express the relationship between synonymous distance ($d_S$) and the mutation distance ($d_{BCS}$) for a given gene $i$ by

$$d_S = (1 - f_{i})d_{BCS},$$

where $f_i$ refers to the fraction of mutations underestimated due to selection against the unpreferred codons and estimation biases. This equation is parallel in concept to that well known for the neutral theory (Kimura 1983), showing that the rate of synonymous substitution becomes equal to the rate of point mutation under the strict neutrality. The synonymous distance for the genes with the smallest (or no) codon usage bias is one of the best candidates of the mutation distance, as there is the least selection or evolutionary distance estimation bias (e.g., Dunn, Bielawski, and Yang 2001). For instance, figure 1a shows the well-known negative relationship between CAI and synonymous distance ($d_{BCS}$) for a given gene ($R^2 = 0.148$ and 0.176 for D. melanogaster and D. simulans, respectively).

![Fig. 1](image.png)

**Fig. 1.**—(a) Relationship of codon adaptation index (CAI) and the number of substitutions per fourfold-degenerate site ($d_S$) between D. melanogaster and D. simulans (62 genes; $R^2 = 0.148$). (b) Relationship between CAI and the base composition skew ($d_{BCS}$) in D. melanogaster genes (146 genes). (c) Relationships between $d_S$ and $d_{BCS}$ between D. melanogaster and D. simulans (62 genes; $R^2 = 0.176$). Fit of the linear regression line is shown in panels a and c. In panel c, higher order regression fit was statistically not significantly better than the linear regression fit shown.
shown). Because genes with the lowest codon usage bias showed a $BCS \approx 0$, the y-intercept in figure 1c could be used as a bias-corrected estimate of the mutation distance.

Based on the observed linear relationship in figure 1c, we can express $f_i$ in equation (1) as a product of $C_i$ and $\eta$ ($f_i = \eta C_i$), where $C_i$ is the $BCS$ for gene $i$ and $\eta$ is the fraction of mutations eliminated per unit $BCS$ for the genome pair compared. We can now rewrite equation (1) to convert the observed synonymous distance ($d_{Si}$) for a given gene into the estimate of number of mutations per site ($d_{ui}$) using the observed base composition skew for that gene ($C_i$).

$$d_{ui} = d_{Si}/(1 - \eta C_i).$$

The value of $\eta$ necessary for using equation (2) is estimated by using all genes for the given species pair. It is obtained by dividing the absolute value of the slope of the linear regression of $BCS$ on $d_{Si}$ by the $y$-intercept of the regression line. For the melanogaster-simulans pair, $\eta = 3.3$. We found that the estimates of $\eta$ for other species pairs with many genes (e.g., melanogaster-yakuba, melanogaster-pseudoobscura, and melanogaster-virilis) were quite similar to that for melanogaster-simulans and statistically not significantly different at the 5% level ($P = 0.74$, 0.76, and 0.48 for melanogaster-yakuba, melanogaster-pseudoobscura, and melanogaster-virilis, respectively). Because the melanogaster-simulans comparison has the largest number of genes, $\eta$ is estimated with the smallest error and is therefore used as a global value for estimating $d_{ui}$ in all further analyses.

**Results**

**Genomic Mutation Clocks**

**Relationship of Mutation Distance and Codon Adaptation**

Figure 2a–d show that the estimated $d_{ui}$ does not depend on the extent of codon adaptation in the analysis of closely as well as distantly related species. The multigene distributions of $d_{ui}$ are symmetrical in nature and have a clear-cut central tendency (fig. 3); means of these distributions provide estimates of genomic mutation distances, $d_{ui}$. This method for estimating genomic mutation distance is useful when building molecular time scales, as it allows for the combining of data from many different genes for different pairs of species because databases are gene rich for some species but poor for most others.

**Relative Rate Tests and Mutation Clocks**

With the ability to compute genomic mutation distances between species, we are in a position to estimate mutation rate and species divergence times as long as mutations accumulate in a clock-like fashion. Therefore, we conducted the relative rate tests (Beverley and Wilson 1984; Wu and Li 1985; Nei and Kumar 2000) of the mutational clock at different levels of evolutionary relatedness. Results from the relative rate tests for five pairs of species are shown in table 1. None of the tests show significant departure from the null hypothesis of mutation rate equality between lineages. Therefore, mutations appear to accumulate at similar rates among diverse Drosophila species, at least up to the melanogaster-pseudoobscura divergence.

**Divergence Time Estimation**

An accurate calibration point is required to anchor the molecular clock to estimate divergence times. This is provided by a well-established divergence time of 5.1 MYA between D. picticornis and other species belonging to the planitibia subgroup based on the time of formation of Kauai in Hawaii (Carson and Clauge 1995). This estimate is considered to be the most robust/reliable estimate and has been used in numerous studies (e.g.,

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**FIG. 2.—Relationship of estimated mutation distances ($d_{ui}$) and codon adaptation index (CAI) for the comparison of D. melanogaster with (a) D. simulans, (b) D. yakuba, (c) D. pseudoobscura, and (d) D. virilis. The correlation between $d_{ui}$ and CAI is not significant in any of the cases ($P > 0.7$, 0.80, 0.3, and 0.4, respectively).**

**FIG. 3.—Multigene mutation distance histogram for comparisons between D. melanogaster and D. simulans (mel-sim), yakuba (mel-yak), pseudoobscura (mel-pse), and virilis (mel-vir). For direct comparison, each multigene distribution was standardized by using the average genomic distance for the corresponding species pair.**
Rowan and Hunt 1991; Takezaki, Rzhetsky, and Nei 1995). With this estimate, the rate of point mutation was computed to be \( \mu = 0.113 \times 10^{-8} \) mutations per site per year per lineage, because the genomic mutation distance between \( D. \) picticornis and other species belonging to the \( planitibia \) subgroup was \( 0.113 \pm 0.0218 \). We used this rate to convert the genomic mutation distance to time for all other species comparisons. The variance of divergence time (\( V_t \)) was estimated by \( V_t = \frac{V_d^2}{\mu^2} + \frac{V_d^2}{\mu^2} \), where \( V_d \) and \( \mu \) represent the variances of \( d_p \) and \( \mu \), respectively.

In the estimation of divergence times of the \( ananassae \) subgroup from the \( melanogaster \) subgroup, we found that the average of uncorrected synonymous distances was much higher than that for the \( obscura-melanogaster \) comparison. This is unexpected because the species in the \( ananassae \) subgroup are considered to be more closely related to \( D. \) melanogaster than those in the \( obscura \) group. In fact, in the \( ananassae-melanogaster \) sequence comparisons, the observed heterogeneity of substitution pattern was higher than that expected in \( >99\% \) Monte-Carlo replicates of the disparity index test (Kumar and Gadagkar 2001) for 7 of 10 genes. For the other 3 genes, this percentage was \( >80\% \). This extreme substitution pattern heterogeneity in the \( ananassae \) lineage might be a reason for the high synonymous distance estimates. However, there were three genes that were shared by species belonging to the \( ananassae \) subgroup, \( melanogaster \) subgroup, and \( obscura \) group. In those genes, the mutation distance for the \( ananassae-melanogaster \) comparison was always lower than that for the \( obscura-melanogaster \) comparison. This allowed us to use those genes to estimate \( ananassae-melanogaster \) divergence time by using the \( obscura \) group as an outgroup. For each gene, we divided the mutation distance for the \( ananassae-melanogaster \) divergence by the mutation distance for the \( obscura-melanogaster \) divergence and multiplied it by the \( obscura-melanogaster \) divergence time to obtain the \( ananassae-melanogaster \) divergence time. Finally, the average divergence time was obtained from three gene-specific estimates to reduce the effect of gene sampling errors. This estimate will need to be refined in the future as more sequences become available.

**Divergence Times in Fruit Fly Evolution**

Figure 4 shows the species divergence times for taxon-pairs with multiple genes (3–64 genes); 2,977 pairwise sequence comparisons from 176 genes were used in this analysis. The estimated molecular time scales for major divergence events are in good accordance with the inferences from biogeography and fossil records. The divergence of \( melanogaster + simulans \) from \( orena + erecta \) and from \( yakuba + teissieri \) (12.6–12.8 MYA) is in agreement with previous inferences based on biogeography (Lachaise et al. 1988). A fossil of a \( Scaptomyza \) species was found in Dominican amber with a minimal age of 23 Myr (Grimaldi 1987). Because \( Scaptomyza \) originated in Hawaii from a common ancestor of the Hawaiian \( Drosophila \) (Tamura et al. 1995; Tatarenkov, Zurovecova, and Ayala 2001), their divergence time should be at least 23 Myr. A molecular estimate of 30.5 (\( \pm 6.6 \)) MYA is consistent with this requirement. In a similar way, the oldest time estimate of 62.9 \( \pm 12.4 \) MYA is consistent with the maximum possible divergence time of 80 Mya between subgenera \( Drosophila \) and \( Sophophora \) as inferred from biogeographic considerations (Beverley and Wilson 1984). These agreements support potential appropriateness of the calibration of the genomic mutation clock.

However, our genomic mutation clock-based estimates are significantly older for some speciation events that have been previously dated by molecular clock methods. In particular, the \( melanogaster-simulans \) divergence based on 62 genes is twice as old as thought before (Thomas and Hunt 1993; Russo, Takezaki, and Nei 1995); this result is statistically significant (\( P < 0.02 \); using a Z-test to compare our estimate with the currently accepted value of 2.5 MYA). Previous studies mainly used \( Adh \) gene data whose sequences are available from the largest number of \( Drosophila \) species. Although we have used the same calibration point (Hawaiian species divergence) as did the other studies, they did not account for codon usage bias differences between the Hawaiian species and other species. The effect of this factor is clearly evident in figure 5a for the \( Adh \) gene data; the rate of uncorrected synonymous substitution for the \( Sophophora \) lineage leading to \( D. \) melanogaster is almost half of that for the subgenus \( Drosophila \) lineage leading to the Hawaiian species. This correlates with the difference in codon usage bias between \( D. \) melanogaster and Hawaiian \( D. \) picticornis, which produces much higher reduction in synonymous distances for the \( Sophophora \) lineage than those for the Hawaiian species and leads to considerable underestimation of the divergence times.
correcting for the effect of the codon usage bias difference on synonymous substitutions using equation (2), the mutation rate difference observed in Adh genes between the two subgenera disappears (fig. 5b). Consequently, our mutation distance for the Adh gene alone dated the melanogaster-simulans and melanogaster-yakuba divergence times to be 4.0 ± 1.7 and 13.7 ± 3.7 MYA, respectively, whereas the uncorrected synonymous distances for fourfold-degenerate sites suggested that they were 2.1 ± 0.9 and 6.8 ± 1.9 MYA. It is interesting to note that our older time estimates based on corrected synonymous distances are in a good agreement with those reported by Beverley and Wilson (1984). Therefore, it is important to use corrected synonymous distance (mutation distance) in building reliable fruit fly evolutionary timescales.

In figure 4, we have also presented three divergence times between very closely related species pairs, i.e., pseudoobscura-persimilis, simulans-mauritiana, and pseudoobscura-miranda. These estimates do not exclude the effect of ancestral population polymorphism, as population polymorphism data do not currently exist for many genes. Therefore, they may be considered to be maximum estimates and will need to be revised in the future.
Single Gene Molecular Clocks

The linear fit of species divergence times for the mutation distances observed for each individual gene, with divergence times estimated by using all the genes, is shown in table 2. For almost all genes shown, $R^2$ values in linear regressions are very high for mutation distances and the average rates of mutation are rather similar. The multigene histogram of these mutation rates is shown in figure 6a. This distribution has the average rate ($\pm 1\ SE$) of 0.0114 $\pm$ 0.0003. These mutation rates are contrasted with the nonsynonymous substitution rates for the same set of genes shown in figure 6a. The L-shaped distribution of the nonsynonymous substitution rates shows that Drosophila genes are under strong purifying selection, with only one gene showing a nonsynonymous substitution rate significantly higher than the average genomic mutation rate. This is the Acp26Aa gene, which is well known to be under strong positive selection (Tsaur and Wu 1997). Some of the other genes showing high nonsynonymous substitution rates were Acp29AB, mei-218, and rux; all of which have already been recognized as rapidly evolving at the protein sequence level (Aguade 1999; Avedisov et al. 2001; Manheim et al. 2002). However, the nonsynonymous rates were much lower than the average genomic mutation rate in all of these cases.

The protein distances for individual genes show much poorer linearity with species divergence times, as compared to that for mutation distances (table 2). Many genes for which amino acid substitutions are known not to follow a clock-like accumulation are clearly exposed. Both Adh and Gpdh show extremely poor fit with species divergence times (Rodriguez-Trelles et al. 2001a), suggesting that their protein evolution is not clock-like. In contrast, some genes such as amd, yellow, and rux appear to accumulate amino acid substitutions in a more clock-like fashion. It is notable that mutation and protein evolutionary rates do not show a significant correlation (fig. 6b), clearly establishing the independence of the estimated mutation and protein evolutionary rates.

Discussion

We have presented a method to correct the bias introduced by the codon adaptation when estimating evolutionary divergence at synonymous sites for inferring mutation distances. Our method works by employing the relationship of the codon usage bias with the synonymous divergence observed in the Drosophila genomes and produces a measure of the average genomic mutation distance. However, the codon usage bias is caused not only by selection but also by mutational bias in the strictly neutral sites. For example, the G+C contents of introns and noncoding regions are about 40% in a variety of Drosophila species, as the direction of mutation is biased toward nucleotides A and T (Moriyama and Hartl 1993; Bergman and Kreitman 2001). This suggests that the expected codon usage in the absence of codon selection is already biased and that $BCS = 0$ (used in our formulation) may not correspond to the case of zero codon adaptation. The violation of the assumption may lead to biased estimation of the mutation distance. However, it is well known from studies of mammalian genomes (e.g., Hughes and Yeager 1997) that the third codon position G+C content is about 10% higher than that observed in introns and intergenic regions. If this is true for Drosophila genomes as well, then we expect about 50% G+C content to be the baseline G+C content for the fourfold-degenerate sites in the Drosophila genomes. This is indeed the case because $>97\%$ genes show fourfold-degenerate site G+C content equal to or higher than 50% in our study. At any rate, it is virtually impossible to estimate the exact base composition at the strictly neutral synonymous sites. Therefore, the average genomic mutation rate reported here should be considered an approximation.

Comparative sequence analyses inherently generate average genomic rates among species. Therefore our estimates are average estimates among species. However, the similarity of base compositions of introns and noncoding regions over widely divergent Drosophila species (Moriyama and Hartl 1993; Bergman and Kreitman 2001) suggests that the mutation bias has not changed.
much during fruit fly evolution and that our estimates may be close approximations for species-specific mutation rates. (See, however, a small departure for D. willistoni reported by Bergman et al. [2002].) This permitted us to use genomic mutation distances from different gene sets from different species pairs to maximally utilize the available data in our analyses.

Relative rate tests conducted using the mutation distances at different levels of taxonomic divergence show that the null hypothesis of the presence of a mutation clock in diverse fruitfly species is not rejected. This allows for assuming a mutation clock and inferring a temporal pattern of species divergences during fruit fly evolution leading to *D. melanogaster*. In fact, the order of divergences for all the species groups (obscura and willistoni) and subgroups (takahashii, montium, and ananassae) belonging to the subgenus *Sophophora* from *D. melanogaster* (fig. 4) is consistent with previous studies for molecular phylogeny of *Sophophora* using multiple genes (Goto and Kimura 2001; O’Grady and Kidwell 2002).

On the basis of the divergence times estimated, it is interesting to speculate about the temporal pattern of speciation events in the evolutionary history leading to *D. melanogaster*. Figure 4 suggests that the speciation events have not occurred regularly in time, as several events are clustered. Three independent pairs of sibling species (*melanogaster-simulans*, yakuba-teissieri, and orena-erecta) diverged within a short time (5.4–6.8 MYA). Evolutionary divergences among these three pairs also occurred in a short span of time 10.4–12.8 MYA. The *melanogaster* subgroup (containing these six species) diverged from the cluster of the three major subgroups in the *melanogaster* group (containing ananassae, montium, and takahashii subgroups) during the Late to Middle Eocene (35–45 MYA). The Hawaiian *Drosophila* also diverged from the lineage leading to *D. viridis* during this time period. Finally, both the *obscura* and *willistoni* groups diverged from the *melanogaster* group about 55 to 62 MYA after the divergence of the subgenera *Drosophila* and *Sophophora*, close to the K-T boundary, 63 MYA. These clustered timings of divergence are compatible with “radiations” proposed by Throckmorton (1975) to explain coincidental distribution patterns of species from independent species groups.

Many of the clustered divergence times either coincide or fall close to the periods of major climate changes during the Cenozoic. Marine sediment records suggest that important cooling steps occurred during the Late Miocene (5.0–6.5 MYA) and the Middle Miocene (12–15 MYA) (Kennett 1995; Zachos et al. 2001) era that coincide with a number of divergence time estimates (fig. 4). By contrast, there is a paucity of speciation events during Oligocene to the Early Miocene, periods with relatively uniform climatic temperatures or warming. An event mapped to this period is the Hawaiian *Drosophila-Scaptomyza* split which occurred in Hawaii, where local volcanic activities are thought to be responsible for speciation (Carson 1992). There are also clear correspondences of older species divergence events with climatic cooling. Although we have many species from most of the major species groups and subgroups related to *D. melanogaster* in our analysis, speciation patterns for independent species groups and subgroups need to be examined with a number of genes to generalize these inferences. Nevertheless, if the observed correspondence between the time of species divergences and paleoclimate changes is true, it supports Wallace’s hypothesis for a rapid species change resulting from climatic change (Wallace 1870a, b). In the present case, the factor is postulated to be climatic cooling in the Cenozoic. A major consequence of this cooling was an extensive increase in aridification in the middle to low latitude regions, which lead to expansions of savannas and grasslands as well as the fragmentation of forests (Kennett 1995) that were primary habitats of ancestral fruit fly species and populations (Throckmorton 1975). The adaptation to the newly arisen dry environment and the allopatry caused by the forest fragmentation are potential causes for stimulating fruit fly speciation. The former adaptation is supported by the distribution patterns of *D. teissieri* and *D. yakuba*, which are adapted to forests and savannas, respectively (Lachaise et al. 1988). The allopatric speciation is also plausible from the overlapping distribution patterns for independent species pairs, say, *D. melanogaster-simulans* and *D. teissieri-yakuba* (Lachaise et al. 1988).
Therefore the mutation clock proposed here provides opportunities to get an important glimpse of speciation processes and mechanisms when they are examined in the context of the contemporary earth history and environmental changes. Although the current view is still mostly speculative, the correlation between speciation events and Cenozoic climatic cooling will become better understood with accumulation of gene sequence data for other fruit fly species. The inferred temporal pattern of speciation from these efforts will also be useful in selecting genomes for sequencing and annotation, calibrating the tempo of DNA loss, building temporal contexts of origin and horizontal transfer events of the transposable elements, and understanding timing of gene duplications, chromosomal changes, and evolution of genome anatomies in general (Petrov and Hartl 1998; Silva and Kidwell 2000; Bergman et al. 2002; Kaminker et al. 2002).

Acknowledgments

We thank Adriana Briscoe, Thomas Dowling, Michael Rosenberg, Martin Wojciechowski, Margaret Kidwell, and Alan Filipski for their comments on an earlier draft of this manuscript. We also thank Balaji Ramanujam for providing invaluable technical assistance. This work was supported by a research grant from the National Institutes of Health, and Burroughs Technology, Japan, to K.T., and from the National Science Foundation, National Institutes of Health, and Burroughs Wellcome Fund, USA, to S.K.

Literature Cited


Thomas Eickbush, Associate Editor
Accepted August 5, 2003