

Analysis of *Drosophila* Species Genome Size and Satellite DNA Content Reveals Significant Differences Among Strains as Well as Between Species

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ABSTRACT

The size of eukaryotic genomes can vary by several orders of magnitude, yet genome size does not correlate with the number of genes nor with the size or complexity of the organism. Although “whole”-genome sequences, such as those now available for 12 *Drosophila* species, provide information about euchromatic DNA content, they cannot give an accurate estimate of genome sizes that include heterochromatin or repetitive DNA content. Moreover, genome sequences typically represent only one strain or isolate of a single species that does not reflect intraspecific variation. To more accurately estimate whole-genome DNA content and compare these estimates to newly assembled genomes, we used flow cytometry to measure the 2C genome values, relative to *Drosophila melanogaster*. We estimated genome sizes for the 12 sequenced *Drosophila* species as well as 91 different strains of 38 species of Drosophilidae. Significant differences in intra- and interspecific 2C genome values exist within the Drosophilidae. Furthermore, by measuring polyploid 16C ovarian follicle cell underreplication we estimated the amount of satellite DNA in each of these species. We found a strong correlation between genome size and amount of satellite underreplication. Addition and loss of heterochromatin satellite repeat elements appear to have made major contributions to the large differences in genome size observed in the Drosophilidae.

THE evolutionary processes associated with the wide spectrum of eukaryotic genome sizes have eluded biologists for decades. The so-called “C-value paradox” refers to our lack of understanding as to how and why there is so much variation in eukaryotic genome size (for reviews see HARTL 2000; PETROV 2001). For example, the mountain grasshopper *Podisma* has an estimated genome size 100-fold that of the fruit fly *Drosophila melanogaster* and ~6-fold larger than the human genome (HARTL 2000; BENSASSON *et al.* 2001; PETROV 2001). Genome size in these examples clearly does not correlate with the number of genes found in each genome or with the complexity of the organism. It appears, instead, that the vast differences in genome size are a result of repetitive DNA sequences that litter eukaryotic genomes in one form or another (HARTL 2000). These observations raise several interesting questions: First, how have genomes of closely related species changed and have repetitive sequences contributed to the evolution of closely related genomes and distantly related species alike? Second, what are the molecular mechanisms through which ge-

nomes change their DNA content? Finally, and most interestingly, are such changes in eukaryotic genome size under selection? The availability of genome sequences, especially of closely related species such as the 12 *Drosophila* genomes, now make it possible to compare whole genomes and address some of these questions.

How have genomes changed? Various models have attempted to describe how genomes have evolved to contain more or less DNA (for reviews see BRITTEN and DAVIDSON 1971; HARTL 2000; PETROV 2001, 2002). Using *Drosophila*, studies attempting to detect global trends in genome size have focused on measurements of transposable elements, pseudogenes, intron, exon, and intergenic lengths (PETROV *et al.* 1996; MORIYAMA *et al.* 1998; PETROV and HARTL 1998). Such studies have been illuminating and suggest that global forces determine the growth and contraction of disparate genomic elements. For example, large genomes tend to have larger intergenic distances, introns, and exons (MOREAU *et al.* 1985). However, repetitive DNA sequences account for the bulk of the vast differences that have been reported (HARTL 2000). For example, the closely related *D. nasutoides* and *D. simulans* have been reported to have 56 and 5% satellite repeat DNA, respectively (ZACHARIAS 1986; LOHE and BRUTLAG 1987).

By what mechanisms have these genomes changed size? Random deletions/insertions, polyploidization, and proliferation of transposable elements are thought

This article is dedicated to Joao Torres Leiva-Neto (1974–2005), who was one of our most enthusiastic and dedicated students and without whom this study would not have been possible.

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to contribute to genome change (for review see HARTL 2000). Also, certain sequences, for example, repetitive elements typical of heterochromatin, may have repeat-specific shrinkage mechanisms, such as unequal meiotic exchange between sister chromatids or replication errors (BRITTEN and KOHNE 1968; SOUTHERN 1975; SMITH 1976; STEPHAN and CHO 1994; PETROV 2001). Understanding the levels and distributions of heterochromatic repetitive elements across a range of related species will aid in discriminating among the potential responsible mechanisms.

Given that most eukaryotic genomes contain vast amounts of repetitive sequences (HARTL 2000), understanding how these sequences contribute to genome evolution is critical. Moreover, it is becoming increasingly clear that heterochromatic repeats and tandem array repeats are not “junk DNA,” but rather serve critical functions, such as meiotic chromosome pairing, epigenetic maintenance of centromere function, and other epigenetic processes (HAWLEY *et al.* 1993; DERNBURG *et al.* 1996; SUN *et al.* 1997; ALLSHIRE 2002; REINHART and BARTEL 2002; CAM *et al.* 2005; CHANDLER 2007). However, the repetitive nature of heterochromatic and other DNAs makes them difficult to clone and sequence (SUN *et al.* 2003). Consequently, assembled genome sequences often do not accurately represent heterochromatic content and thus underestimate total genome size as well as repeat sequence content.

Genome size estimates are available for 70 species of the family Drosophilidae (POWELL 1997; ASHBURNER *et al.* 2005; <http://www.genomesize.com>) and clearly exhibit large differences among and within species. Multiple estimates exist for several species and suggest intraspecific genome size differences of up to 50% for some. In strains of *D. melanogaster*, the intraspecific genome size variation was attributed to differences in heterochromatin content (HALFER 1981). Scant information is available, however, regarding the heterochromatin satellite DNA content of many other species, and thus available genome size estimates have limited usefulness in addressing evolutionary questions. The majority of existing estimates are from unpublished studies and thus details regarding the methodology, tissues, and strains used cannot be ascertained. Remaining estimates were performed with a range of different techniques, such as flow cytometry, Feulgen densitometry, molecular weight determinations, and sequencing, and employed different tissue types such as ovaries, sperm, testes, brains, whole bodies, and hemocytes. These methodological inconsistencies, coupled with an absence of information on the contribution of various repeat sequences to the observed genome size variability, necessitate a new approach that will provide accurate simultaneous measures of both genome size and satellite DNA content across the Drosophilidae. Of special interest are those 12 species for which whole-genome sequences are now available (<http://rana.lbl.gov/drosophila/>).

In this study, we address the following questions: (1) What is the range of genome sizes across the Drosophilidae?, (2) What is the range of variation within species for genome size?, and (3) What is the contribution of heterochromatic satellite DNA to intra- and interspecific variability in genome size? To address these questions, we ascertained the genome sizes of 91 strains from 38 species within the Drosophilidae, including the 12 sequenced species (<http://rana.lbl.gov/drosophila/>). Using flow cytometry, we determined the genome sizes and the fraction of each of these genomes that is under-replicated in ovarian follicle cells. Although follicle cells from all 38 species terminate with 16 complement (16C) ploidy, we observed dramatic differences in the fraction of the 2 complement (2C) genome that is actually replicated in each species. This indicates measurable differences in under-replicated satellite content. We also found a strong correlation between genome size and amount of satellite DNA, suggesting that variation in heterochromatic DNA contributes significantly to genome size evolution in the Drosophilidae.

MATERIALS AND METHODS

Species and strains used: To identify potential strain differences, we examined more than one strain of each species—a total of 91 different strains from 38 species. All strains and species are available for future analysis and most are banked in the Tucson *Drosophila* Species Stock Center and the Bloomington (Bl) *Drosophila* Stock Center (supplemental Table 3 at <http://www.genetics.org/supplemental/>). One strain (H2AvD-GFP; CLARKSON and SAINT 1999) and one *D. virilis* strain (no. 2465, origin unknown but likely from M. Pardue, Massachusetts Institute of Technology) are available upon request from G. Bosco. Since Bloomington stock numbers can change over time, genotypes for each *D. melanogaster* strain are shown in supplemental Table 3 at <http://www.genetics.org/supplemental/>.

Preparation of nuclei and flow cytometry: We dissected 10–20 ovary pairs in Grace's insect medium (GIBCO, Grand Island, NY) and placed them into 1.7-ml tubes with 0.8 ml of medium. Grace's medium was removed and 700 μ l filtered ice-cold PARTEC buffer (200 mM Tris-HCl pH 7.4, 4 mM MgCl₂, 0.1% Triton X-100) was added to the 1.7-ml tube with the ovaries and then placed into a 60-mm petri dish and homogenized with a single-edged razor blade. Chopped ovaries were filtered twice over cheesecloth (\sim 3 cm²) and once through a 30- μ m mesh (Sefar) and collected in a flow cytometry tube (Sarstedt). Another 700 μ l of PARTEC buffer was used to wash the petri dish, filtered, and pooled into flow cytometry tubes.

Two nucleic-acid-binding fluorescent dyes were used, propidium iodide (PI) and 4',6-diamidino-2-phenylindole (DAPI). For DAPI staining, nuclei in tubes were placed on ice and 20 μ l of DAPI (100 μ g/ml) were added. Samples were analyzed on a PARTEC CCA-II flow cytometry machine (PARTEC). For PI staining, we used the same protocol as above with the addition of 50 μ l RNase A (1 mg/ml) and 100 μ l PI (1 mg/ml) to each sample. PI measurements were done on a FACScan flow cytometer (Becton Dickinson) at several thousand nuclei per second.

For both DAPI and PI measurements, each sample was compared to a *D. melanogaster* control (*y^{1w}* Bloomington no. 1495, hereafter referred to as *D.m. yw*) that was prepared at the same time for each sample. Both PARTEC CCA-II and FACScan machines were calibrated to flow rates and gain settings for the

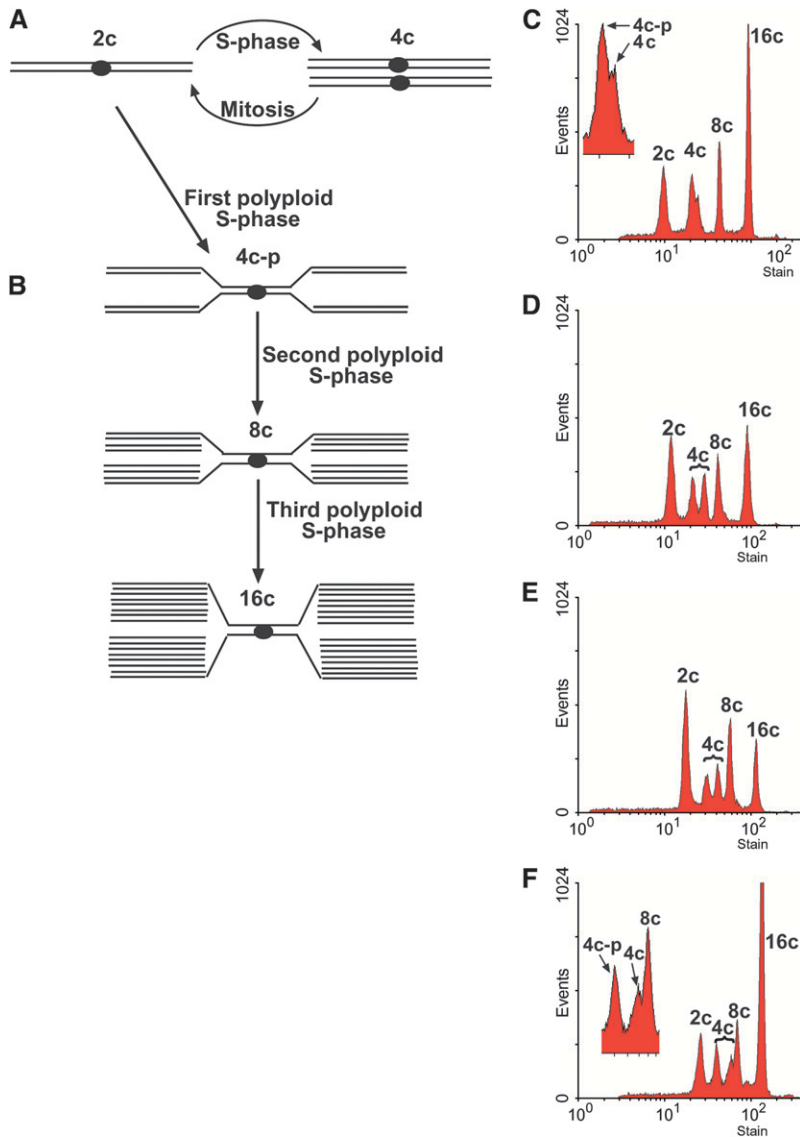


FIGURE 1.—*Drosophila* polyplod follicle cells underreplicate satellite DNA repeats. Proliferating follicle cells duplicate their entire genomes and cycle from 2C to 4C and after mitotic division back to 2C (A). 2C cells enter their polyplod state by replicating their euchromatic sequences and replicate little or no centric/pericentric satellite repeat sequences (B). Consequently, 4c-p cells have less 4C DNA content, and a second and third round of polyplod S-phases produce 16C cells with vastly underreplicated satellite DNA. Flow cytometry histograms of follicle cell nuclei from (C) *D. melanogaster*, (D) *D. grimshawi*, (E) *D. immigrans*, and (F) *D. virilis* are shown by illustrating the four major 2C, 4C, 8C, and 16C ploidy peaks where the x-axis represents arbitrary fluorescent units and the y-axis is the number of nuclei. Note that the 4C peak can be resolved into two peaks (see insets in C and F), where the 4C peak from mitotic proliferating cells has more DNA content than the 4C-p peak. This is because follicle cells undergoing polyplodization fail to replicate the centric and pericentric satellite repeats and thus have less DNA than mitotic 4C cells, as described in A. In larger genomes such as (D) *D. grimshawi*, (E) *D. immigrans* and (F) *D. virilis*, the extent of underreplication can be seen by a dramatic shift of all polyplod peaks to the left. The most extreme example is seen in (F) *D. virilis* where the 8C peak nearly overlaps the normal mitotic cell 4C peak (see inset), suggesting that about half of the genome fails to replicate. This is consistent with measurements of ~48% heterochromatin content in *D. virilis* (see Table 5). We observed underreplication in all 91 strains from all 38 species that we examined.

D. m. yw control. In all cases, a minimum of three biological replicates was performed on each strain, and a minimum of 10^4 nuclei was measured for each replicate.

Determination of flow cytometry values and statistical analysis: Histograms exhibiting four peaks (2C, 4C, 8C, and 16C) were obtained for polyplod follicle cells (Figure 1). The mean fluorescence intensity for each peak was obtained and this fluorescence value is proportional to DNA content as previously described for follicle cell nuclei (LILLY and SPRADLING 1996; LEACH *et al.* 2000; BOSCO *et al.* 2001). As ANOVA revealed no significant differences among replicates for a given strain, they were averaged (data not shown and Table 1). This average fluorescent intensity was divided by its *D. m. yw* control, yielding a normalized estimate of 2C DNA content, relative to *D. m. yw*. For each of the three biological replicates for each strain, 16C/2C ratios were determined and then averaged to obtain an average 16C/2C ratio for each strain.

Conversion of 2C values to picograms and megabases: To convert relative genome sizes to megabase values, we produced a best-fit regression line for experimentally measured 2C flow cytometry values and the corresponding published genome sizes for *D. melanogaster* and *D. virilis*. (LAIRD 1971, 1973; RASCH *et al.* 1971; KAVENOFF and ZIMM 1973; MULLIGAN and RASCH

1980; CELNIKER *et al.* 2002; HOSKINS *et al.* 2002; BENNETT *et al.* 2003). Two best-fit curves (one for PI and another for DAPI) were obtained, which then were used to convert 2C measurements into megabase values. The advantage of this method is that it takes into account complex relationships between 2C flow cytometry values and DNA content for different species. One disadvantage is the lack of information on the *D. virilis* strains used previously for genome size estimates. Consequently, we used an average from two different studies (KAVENOFF and ZIMM 1973; LAIRD 1973) and must assume that these *D. virilis* strains are sufficiently close to the five strains examined in this study.

Relative 2C values used for conversion to megabases are shown in supplemental Tables 1 and 2 at <http://www.genetics.org/supplemental/>. DAPI relative 2C values were first corrected for A:T bias as described below and in Figure 2A. Picograms were calculated from megabases based on the conversion $0.1 \text{ pg} = 97.8 \text{ Mb}$.

Estimates of underreplicated satellite content: The expected DNA content of 16C polyplod follicle cells is eight times the raw 2C value ($8 \times 2C$). Observed raw 16C values obtained from PI flow cytometry are less than the expected values because heterochromatic sequences do not replicate completely if at all in

follicle cells (Figure 1) (GALL *et al.* 1971; HAMMOND and LAIRD 1985a; LILLY and SPRADLING 1996; LEACH *et al.* 2000). Thus, the difference between the expected and the observed 16C values reflects the fraction of each genome that is underreplicating satellite repeats $[(8 \times 2C) - 16C]$. For values obtained by PI fluorescence, the following formula was used to calculate the percentage of underreplication in 16C follicle cells: $[(8 \times 2C) - 16C] / (8 \times 2C) \times 100$. The percentage of underreplication is an estimate for the heterochromatic satellite DNA content in each genome.

Determination of the expected 16C ploidy DNA contents (*i.e.*, $8 \times 2C$) with DAPI data is confounded by the fact that DAPI values are skewed by A:T content, and therefore 2C values and 16C values reflect DNA content plus A:T richness. Consequently, estimates of underreplication determined by DAPI will be less precise than those derived from PI measurements, and DAPI values must first be normalized for the A:T bias. To normalize DAPI 16C/2C values, we used the following formula: normalized DAPI 16C/2C = $[(PI\ 16C/2C\ D.m.\ yw) / (DAPI\ 16C/2C\ D.m.\ yw)] \times DAPI\ 16C/2C$ for each strain. Normalized DAPI percentages of underreplication values were determined by multiplying the normalized DAPI 16C/2C by 26%. Because we determined a mean 26% underreplication for four *D. melanogaster* strains by using PI (Table 4), the mean 26% value was used to convert 16C/2C values that were normalized to *D. melanogaster*.

Chromocenter measurements and immunofluorescence: Ovaries were dissected and prepared for DAPI (0.05 μ g/ml final) and immunofluorescence (HARTL *et al.* 2007). Rabbit antidimethyl lysine-9 on histone H3 (Upstate) was used at 1:100 dilution and visualized with Cy3-goat anti-rabbit (Jackson ImmunoResearch) at 1:250 dilution. Stage 13 follicle cell nuclei were imaged with a Nikon Eclipse E800 microscope and a $\times 40$ objective using a RT Monochrome SPOT Model 2.1.1 camera. All settings were kept identical for all samples although background signal varied among samples. Nuclear and chromocenter areas were determined with the Adobe Photoshop 7.0 Polygonal Lasso tool, and the total areas for each nucleus and chromocenter were determined in pixels using the Image histogram function. The area of the chromocenter, as determined by DAPI and histone H3 dimethyl-lysine-9, was normalized to the total nuclear area. An average normalized chromocenter area for each species was calculated. For each of the three species examined, 35 different cells were measured. Standard errors and *P*-values using a two-tailed test were determined using MS Excel.

RESULTS

Fluorescent flow cytometry can accurately estimate genome size: As genome size estimates were previously available for *D. melanogaster* (LAIRD 1971; RASCH *et al.* 1971; KAVENOFF and ZIMM 1973; MULLIGAN and RASCH 1980; CELNIKER *et al.* 2002; BENNETT *et al.* 2003) and *D. virilis* (KAVENOFF and ZIMM 1973; LAIRD 1973), we assessed the ability of PI and DAPI flow cytometry to accurately reproduce the previously described genome size differences for these two species. For example, previous estimates described the *D. virilis* genome to be much larger than *D. melanogaster* and to have a higher heterochromatin content (GALL *et al.* 1971; SCHWEBER 1974). We conducted a set of preliminary studies on multiple strains of *D. melanogaster* and of *D. virilis* and determined the fluorescence intensity for follicle cell nuclei with 2C and 16C ploidy, relative to *D. melanogaster yw* controls

(Table 1). We performed flow cytometry using PI fluorescence for four *D. melanogaster* and four *D. virilis* strains. Using PI as the dye, ANOVA detected significant species differences, but not strain or replicate differences in 2C values or 16C/2C values (Table 2A). ANOVA performed on measurements of the same 4 plus 6 additional *D. melanogaster* strains (10 total) and on the same 4 plus 1 additional (5 total) *D. virilis* strains with DAPI revealed significant species and strain, but not replicate, differences (Table 2B). We conclude that both dyes detect interspecific genome size differences. Comparison of the DAPI 2C values for each of the *D. virilis* strains to *D.m. yw* revealed a 2.25- to 2.71-fold difference. For PI 2C values, there was a 1.7- to 2.09-fold difference between *D. virilis* and *D.m. yw* (Table 1). Our 2C values fit very well with values for *D. virilis* genome sizes previously estimated to be 1.75- to 2.26-fold larger than *D. melanogaster* (KAVENOFF and ZIMM 1973; LAIRD 1973; J. Spencer Johnston as referenced in Table 5.2 of ASHBURNER *et al.* 2005). This and previously published work demonstrate that flow cytometry provides a valid method for determining genome size when an appropriate control is used (JOHNSTON *et al.* 1999; BENNETT *et al.* 2003).

Effects of dye on genome size measures: In general, 2C DAPI values for most strains, relative to *D. melanogaster*, were elevated when compared to 2C values obtained by PI (Table 1 and supplemental Tables 1 and 2 at <http://www.genetic.org/supplemental/>). DAPI binding preference for A:T sequences has been physically documented (WILSON 1990; COLSON *et al.* 1995, 1996), and its preferential fluorescence for A:T-rich DNA in flow cytometry also has been described (JOHNSTON *et al.* 1999; MEISTER 2005). Moreover, cytological changes in DAPI fluorescent intensity accurately correlate with physical changes in A:T-rich repeat content in *D. melanogaster* polyploid cells (LILLY and SPRADLING 1996; ROYZMAN *et al.* 2002). Discrepancies between DAPI and PI 2C values therefore suggest that most, but not all, species have A:T-rich genomes.

Given the A:T content bias, we plotted DAPI 2C values against PI 2C values to assess whether only some or most species exhibit a DAPI bias (Figure 2A). If DAPI and PI values are equivalent, we would expect a linear relationship with a slope approximately equal to 1. Interestingly, although DAPI values increased with PI values, DAPI values increased at a greater rate (Figure 2A). The trend was highly significant ($P < 0.0001$), consistent with larger genomes having more A:T-rich satellite DNA, thus leading to an exaggerated DAPI signal. A simple conversion from DAPI-derived values to picograms or megabases thus was not possible, especially for A:T-rich genomes, without first performing a correction. A linear regression predicts that PI (*y*) values change in relation to DAPI (*x*) values as described by the equation $y = 0.3832x + 0.6051$ (Figure 2A). We employed a DAPI correction factor that allowed us to account for A:T bias in DAPI fluorescence values where the corrected DAPI 2C value = $0.3832(\text{observed DAPI 2C}) + 0.6051$. The linear regression shown

TABLE 1
Fold difference for multiple *D. melanogaster* and *D. virilis* strains

Strain no.	PI		DAPI	
	2C \pm SE	16C/2C \pm SE	2C \pm SE (A:T corrected)	16C/2C \pm SE
<i>D. melanogaster</i>				
Bl 2057	1.01 \pm 0.09	6.15 \pm 0.29	1.67 \pm 0.01 (1.25)	6.06 \pm 0.04
Bl 1495	1.00 \pm 0	6.30 \pm 0.03	1.00 \pm 0.07 (0.99)	9.31 \pm 0.13
Bl 4455	0.99 \pm 0.01	6.09 \pm 0.01	1.19 \pm 0.05 (1.06)	8.83 \pm 0.10
Bl 6599	1.32 \pm 0.03	5.49 \pm 0.17	1.29 \pm 0.05 (1.10)	9.17 \pm 0.17
Bl 1785			0.97 \pm 0.01 (0.98)	10.65 \pm 0.07
Bl 576			1.13 \pm 0.08 (1.04)	9.06 \pm 0.08
Bl 1633			1.08 \pm 0.06 (1.02)	8.85 \pm 0.03
H2AvD-GFP			1.11 \pm 0.02 (1.03)	9.09 \pm 0.10
Bl 4269			1.12 \pm 0.04 (1.03)	9.01 \pm 0.05
Bl 189			1.04 \pm 0.11 (1.00)	9.00 \pm 0.14
<i>D. virilis</i>				
15010-1051.00	1.97 \pm 0.03	4.44 \pm 0.07	2.71 \pm 0.01 (1.64)	6.12 \pm 0.03
15010-1051.45	2.09 \pm 0.04	4.73 \pm 0.04	2.54 \pm 0.03 (1.58)	6.08 \pm 0.02
15010-1051.46			2.25 \pm 0.08 (1.47)	5.87 \pm 0.01
15010-1051.87	1.78 \pm 0.06	4.57 \pm 0.02	2.38 \pm 0.15 (1.52)	5.86 \pm 0.03
2465	1.70 \pm 0.30	5.40 \pm 1.16	2.34 \pm 0.11 (1.50)	5.21 \pm 0.02

2C and 16C/2C values were obtained for multiple strains of *D. melanogaster* and *D. virilis* using either PI or DAPI dyes in flow cytometric measures of the genome size of ovarian follicle cell nuclei. All values represent averages of three biological replicates, except for *D. melanogaster* Bl 1495 and Bl 2057, which were measured in four and six biological replicates, respectively. Standard error (\pm SE) is shown for each value. DAPI values corrected for A:T bias fluorescence as described in Figure 2A and in the MATERIALS AND METHODS are shown in parentheses (A:T corrected). Note that, before bias correction, the DAPI values for *D. virilis* are much higher than the PI 2C values whereas this dye effect is minimal in *D. melanogaster* 2C values. This reflects a greater total A:T content in *D. virilis*.

in Figure 2A was then utilized to determine the A:T-content-corrected DAPI 2C values (Table 1, A:T corrected). These corrected values were then used to determine genome sizes (Table 3).

TABLE 2
ANOVA analysis of species and strains

	Mean square	Sum-of-squares	F-ratio	P-value
A. PI: <i>D. melanogaster</i> and <i>D. virilis</i>				
2C/2C				
Species	2.88491	2.8849099	53.9511	<0.001
Strain	0.021983	0.0659505	0.0975	0.9604
Replicate	0.029462	0.0589232	0.1380	0.8720
16C/2C				
Species	7.66103	7.661029	12.7873	0.0020
Strain	0.21726	0.651793	0.2008	0.8944
Replicate	0.582162	1.164325	0.5861	0.5668
B. DAPI: <i>D. melanogaster</i> and <i>D. virilis</i>				
2C/2C				
Species	16.4818	16.481775	362.8012	<0.001
Strain	0.709896	6.389066	2.0626	0.0610
Replicate	0.018108	0.036216	0.0413	0.9565
16C/2C				
Species	94.6818	94.68176	111.4805	<0.001
Strain	5.52161	49.69453	2.3710	0.0327
Replicate	0.01223	0.02446	0.0039	0.9961

Total A:T content is positively correlated with genome size: The relative A:T/G:C content of different species can be estimated from the 2C DAPI/2C PI ratio (MEISTER 2005). We took advantage of this DAPI bias to ask how A:T content varies among these *Drosophila* species and whether A:T content was correlated to genome size as suggested by the trend in Figure 2A. Of 48 strains tested (from 30 different species), 33 had log DAPI/PI values greater than zero, indicating that most genomes are A:T rich (Figure 2B, supplemental Table 4 at <http://www.genetics.org/supplemental/>). We observed that, although some smaller genomes are A:T rich, the largest genomes (>250 Mb) are the most A:T rich. Fourteen strains exhibited log DAPI/PI values less than zero, indicating a relatively high G:C content. Interestingly, these G:C-rich genomes were almost exclusively the smallest genomes (<200 Mb), consisting of multiple strains of *D. persimilis*, *D. pseudoobscura*, *D. simulans*, and *D. erecta* (Figure 2B, supplemental Table 4 at <http://www.genetics.org/supplemental/>). This was most pronounced in *D. persimilis* and *D. pseudoobscura*. Consequently, DAPI measurements may underestimate sizes of these genomes and are expected to be lower than PI-derived values, which is in fact what we observed (Table 3).

Genome size estimates: After establishing the efficacy of flow cytometry measurements of 2C *Drosophila* follicle

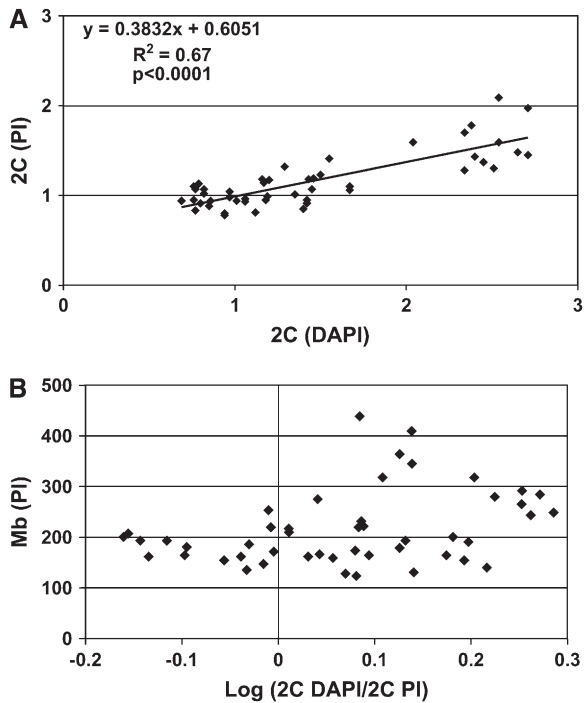


FIGURE 2.—DAPI measurements overestimate DNA content. (A) 2C values relative to *D.m. yw* control for DAPI (*x*-axis) were plotted against their corresponding 2C values for PI (*y*-axis). A trend line was fit to ascertain how DAPI values change relative to PI values. A slope that is < 1 shows that DAPI values increase at a greater rate than PI values. This indicates that as genomes become larger DAPI overestimates DNA content (see text for details) and thus must be corrected. A two-tailed *P*-value was calculated from the correlation coefficient (*R*) and 45 degrees of freedom (d.f.) using Graphpad software. (B) DAPI fluorescence has a A:T bias whereas PI does not. The 2C DAPI/2C PI ratio values for each strain reflect the overall A:T/G:C content of each genome. The log (2C DAPI/2C PI) values (*x*-axis) and the corresponding haploid genome size (*y*-axis) values, as determined by PI 2C, are shown. Note that these measurements are for total genomic A:T/G:C content and may differ substantially from estimates of euchromatic A:T/G:C sequence content.

cells for predicting genome size, we then estimated genome sizes for 91 strains from 38 different species of Drosophilidae. For some species, only 1 strain was available, while for others as many as 10 were tested. Values obtained for individual strains are available in supplemental Tables 1 and 2 at <http://www.genetics.org/supplemental/>. All 38 species were measured with DAPI and 21 also were measured with PI (Table 3). Using PI, the smallest genomes were seen in *D. mercatorum*, *D. mojavensis*, and *D. erecta* while *D. virilis* had the largest. While this pattern was also seen with DAPI, *Chymomyza pararufithorax*'s and *C. rufithorax*'s genomes were slightly larger than that of *D. virilis*.

Follicle cell underreplication is inversely proportional to genome size in all species: We took advantage of the fact that *D. melanogaster* follicle cells that normally become polyploid and have 16C do not completely rep-

licate the centric- and peri-centric heterochromatic satellite DNA (GALL *et al.* 1971; HAMMOND and LAIRD 1985a,b; LILLY and SPRADLING 1996; LEACH *et al.* 2000). Follicle cells undergo three rounds of endoreduplication and terminate with 16C ploidy, as indicated by four major peaks when nuclei are analyzed by fluorescence flow cytometry (Figure 1A). Using this method, we determined that in all 91 strains of 38 species follicle cells terminate DNA replication with 16C ploidy (Figure 1; supplemental Tables 1 and 2 at <http://www.genetics.org/supplemental/>; data not shown). Regulation of follicle cell ploidy thus is a well-conserved developmental process.

An additional 4C-polyploid (4C-p, Figure 1) peak is also evident in the flow cytometry histograms because the majority of the heterochromatic sequences are not completely replicated as these cells progress from 2C through their first polyploid S-phase, resulting in a 4C-p content that has less DNA than a 4C cell undergoing mitosis (HAMMOND and LAIRD 1985a; LILLY and SPRADLING 1996; LEACH *et al.* 2000). The 4C-p peak, since it emits less fluorescence, is always shifted to the left, relative to the 4C peak (Figure 1). The extreme example is satellite repeats that have been estimated to remain at their 2C copy number as polyploidy ensues (GALL *et al.* 1971; LILLY and SPRADLING 1996; LEACH *et al.* 2000). For some species, where extensive underreplication occurs, a distinct additional 4C-p peak is evident (Figure 1, C and F, insets). Moreover, the 16C peaks from different species such as *D. melanogaster* and *D. virilis* are only slightly shifted from one another (compare Figure 1, C and F), indicating that the actual 16C DNA content of these species is not that different. This similarity is observed despite the fact that *D. virilis* 2C DNA content is ~ 1.8 -fold greater than that of *D. melanogaster* (Tables 1 and 4).

We found that the mean PI fluorescence ratio of the 16C/2C values is always < 8 (Table 1; Figure 3A; supplemental Table 1 at <http://www.genetics.org/supplemental/>). The 16C/2C ratio indicates that portion of a given genome that does not fully replicate in follicle cells. Species with larger genomes are expected to have more heterochromatic repeats and therefore replicate a smaller fraction of their 2C genomes in their follicle cells. Indeed, when we plot 2C and 16C/2C values, a clear trend is revealed where larger genomes have smaller 16C/2C ratios (Figure 3A). When we plot DAPI 16C/2C ratios against their corresponding 2C values, we also see a clear negative correlation (Figure 3B; supplemental Tables 1 and 2 at <http://www.genetics.org/supplemental/>). Taken together, these data indicate that *Drosophila* species other than *D. melanogaster* also underreplicate their satellite sequences in follicle cells.

Estimates for underreplicating the percentage of satellite DNA: Although 20% of the *D. melanogaster* genome is estimated to be satellite sequence (LOHE and BRUTLAG 1986), cytological methods and recent heterochromatin sequencing efforts place the heterochromatin content at $\sim 33\%$ (GATTI *et al.* 1976; HOSKINS *et al.*

TABLE 3
Mean genome size and range

Species	Mean \pm SE (<i>n</i>) PI (Mb)	Range of PI	Mean \pm SE (<i>n</i>) DAPI (Mb)	Range of DAPI	Previous estimate (Mb)
<i>C. pararufithorax</i>	284 \pm 6 (1)	—	429 \pm 6 (1)	—	—
<i>C. procnemis</i>	318 \pm 6 (1)	—	260 \pm 7 (1)	—	—
<i>C. rufithorax</i>	292 \pm 6 (1)	—	420 \pm 13 (1)	—	—
<i>D. acutilabella</i>			172 \pm 4 (2)	168–176	—
<i>D. americana</i>	275 \pm 4 (1)	—	240 \pm 14 (2)	226–254	328 ^a (MW)
<i>D. ananassae</i>	215 \pm 5 (3)	210–217	198 \pm 2 (3)	195–202	205 ^b (CY)
<i>D. buskii</i>			194 \pm 5 (1)	—	144 ^b (CY)
<i>D. equinoxialis</i>			304 \pm 9 (2)	295–313	248 ^b (CY)
<i>D. erecta</i>	145 \pm 10 (2)	135–154	139 \pm 2 (2)	137–141	159 ^b (CY)
<i>D. funebris</i>			330 \pm 22 (1)	—	269 ^c (KI)
<i>D. grimshawi</i>	231 \pm 5 (1)	—	247 \pm 13 (1)	—	247 ^d (FD)
<i>D. guttifera</i>	160 \pm 21 (2)	140–181	188 \pm 44 (2)	144–232	—
<i>D. hydei</i>	164 \pm 16 (1)	—	177 \pm 22 (2)	155–199	197–246 ^e (CY, KI, MW)
<i>D. immigrans</i>	299 \pm 19 (2)	279–318	347 \pm 18 (3)	328–382	—
<i>D. littoralis</i>			238 \pm 5 (1)	—	—
<i>D. melanogaster</i>	201 \pm 16 (4)	174–253	195 \pm 10 (10)	167–272	176–180 ^f (CY)
<i>D. mercatorum</i>	128 \pm 5 (1)	—	166 \pm 4 (2)	162–170	—
<i>D. mimica</i>	257 \pm 6 (4)	243–270	387 \pm 8 (3)	373–399	—
<i>D. mojavensis</i>	152 \pm 11 (3)	130–166	183 \pm 3 (3)	180–189	215 ^g (BC)
<i>D. nanoptera</i>			236 \pm 35 (3)	173–295	—
<i>D. novamexicana</i>			244 \pm 20 (2)	224–265	—
<i>D. persimilis</i>	183 \pm 10 (3)	164–193	170 \pm 34 (3)	135–239	197 ^b (CY)
<i>D. pseudoobscura</i>	185 \pm 12 (3)	162–200	135 \pm 6 (3)	125–144	168 ^b (CY)
<i>D. repleta</i>			167 \pm 13 (3)	153–192	—
<i>D. sechellia</i>	166 \pm 5 (2)	162–171	170 \pm 3 (2)	167–173	167 ^b (CY)
<i>D. simulans</i>	160 \pm 11 (6)	123–207	170 \pm 18 (7)	119–235	139–153 ^{h,c} (CY, KI)
<i>D. virilis</i>	404 \pm 21 (4)	364–438	389 \pm 12 (5)	373–429	307–394 ^{h,h} (CY, MW)
<i>D. willistoni</i>	206 \pm 14 (3)	178–222	234 \pm 5 (3)	224–241	235 ⁱ (UN)
<i>D. yakuba</i>	188 \pm 2 (2)	186–190	220 \pm 53 (2)	167–272	173 ^b (CY)
<i>Hirtodrosophila duncani</i>			333 \pm 9 (1)	—	—
<i>S. latifasciaeformis</i>			313 \pm 27 (2)	286–340	195 ^b (CY)
<i>S. lebanonensis</i>			259 \pm 2 (2)	257–260	210 ^b (CY)
<i>S. palmae</i>			168 \pm 9 (1)	—	—
<i>S. stonei</i>			300 \pm 11 (2)	289–311	207 ^b (CY)
<i>Zaprionus badyi</i>			253 \pm 6 (1)	—	—
<i>Z. ghesquerei</i>			153 \pm 7 (1)	—	—
<i>Z. sepsoides</i>			352 \pm 71 (2)	281–423	—
<i>Z. tuberculatus</i>			299 \pm 74 (3)	247–384	—

Mean values for PI and corrected DAPI measurements are for haploid genome size and are from this study. Standard error (SE), the range (lowest and highest values), and the number of strains for each species (*n*) are shown. See supplemental data at <http://www.genetics.org/supplemental/> for specific strain values. DAPI values for larger genomes tend to be less accurate than PI values (see text). For comparison, previously reported genome size estimates are listed in the right-most column. Methods used for determining previous estimates are biochemical analysis (BC), cytometry (CY), kinetics (KI), Feulgen densitometry (FD), and molecular weight (MW), or method unknown (UN).

^aFrom KAVENOFF and ZIMM (1973).

^bFrom J. Spencer Johnston as quoted in Table 5.2 of ASHBURNER *et al.* (2005).

^cFrom LAIRD and MCCARTHY (1969).

^dFrom RASCH (1985).

^eFrom MULDER *et al.* (1968), DICKSON *et al.* (1971), KAVENOFF and ZIMM (1973), and LAIRD (1973).

^fLAIRD (1971), RASCH *et al.* (1971), CELNIKER *et al.* (2002), and BENNETT *et al.* (2003).

^gFrom SCHULZE and LEE (1986).

^hFrom KAVENOFF and ZIMM (1973) and LAIRD (1971).

ⁱFrom POWELL (1997).

2002). By using the 16C/2C ratio we were able to estimate the genomic fraction of each genome that is underreplicated. PI values for 16C/2C indicate underreplication of ~20–31% of the *D. melanogaster* genome in 16C fol-

licle cells while DAPI values show 23–40% (Table 5). These values are surprisingly close to those reported for *D. melanogaster* satellite DNA and heterochromatin content (GATTI *et al.* 1976; HOSKINS *et al.* 2002).

TABLE 4
Genome size and predicted percentage of satellite DNA

Species strain no.	pg \pm SE	Mb \pm SE	Assembly size ^a	% satellite DNA
<i>D. sechellia</i> 14021-0248.25	0.17 \pm 0.004	171 \pm 4	167	24 \pm 0
<i>D. simulans</i> 14021-0251.195	0.17 \pm 0.002	162 \pm 2	142	17 \pm 1
<i>D. melanogaster</i> 14021-0231.36	0.20 \pm 0.017	200 \pm 18	130	24 \pm 3
<i>D. yakuba</i> 14021-0261.01	0.19 \pm 0.011	190 \pm 11	169	23 \pm 2
<i>D. erecta</i> 14021-0224.01	0.14 \pm 0.004	135 \pm 4	153	9 \pm 2
<i>D. ananassae</i> 14024-0371.13	0.22 \pm 0.009	217 \pm 9	231	23 \pm 2
<i>D. pseudoobscura</i> 14011-0121.94	0.20 \pm 0.004	193 \pm 4	153	14 \pm 4
<i>D. persimilis</i> 14011-0111.49	0.20 \pm 0.005	193 \pm 5	188	14 \pm 1
<i>D. willistoni</i> 14030-0811.24	0.23 \pm 0.008	222 \pm 7	237	12 \pm 1
<i>D. virilis</i> 15010-1051.87	0.37 \pm 0.013	364 \pm 13	206	44 \pm 1
<i>D. mojavensis</i> 15081-1352.22	0.13 \pm 0.0	130 \pm 0	194	2 \pm 1
<i>D. grimshawi</i> 15287-2541.00	0.24 \pm 0.005	231 \pm 5	200	32 \pm 0.4

Predicted genome sizes for the 12 sequenced *Drosophila* species. Values, in picograms and megabases, and standard error (\pm SE) for each strain from propidium iodide flow cytometry measurements are shown. The predicted percentage and standard error (\pm SE) of underreplicated heterochromatic satellite DNA is shown for each specific strain.

^aFor comparison, the total assembled sequenced genomes in megabases are shown (<http://insects.eugenesis.org/species/data>). All percentage of satellite DNA estimates are from this study.

Since *D. virilis* has one of the largest genomes (Table 3), we expected this species to have the largest underreplicated DNA content. PI values for 16C/2C *D. virilis* indicate 40–48% underreplication while DAPI values suggest 40–46% (Table 5). These values fit very well with those previously described for *D. virilis* heterochromatin content of 40–42% (GALL *et al.* 1971; SCHWEBER 1974).

To further confirm that underreplication estimates correlate with heterochromatin satellite DNA content, we stained follicle cells with two heterochromatin markers. Centric and pericentric heterochromatin aggregate into a chromocenter in these cells. Chromocenter size and DAPI staining intensity have been found to reflect satellite DNA content (LILLY and SPRADLING 1996; ROYZMAN *et al.* 2002). As shown in Figure 4A, DAPI-stained *D. melanogaster* follicle cell nuclei exhibit bright subnuclear chromocenters. *C. parvifithorax* chromocenters are smaller than in *D. melanogaster*, whereas those of *D. virilis* are larger (Figure 4, D and G). When we used antidimethyl-histone H3 lysine-9 (dmH3-K9) antibodies that recognize methylated H3-K9, a heterochromatin-specific histone modification (ALLSHIRE 2002), the same pattern was observed for these three species (Figure 4, B, E, and H). Species differences in chromocenter size are highly significant (Figure 4J), which is congruent with flow cytometry estimates of 18, 26, and 44% satellite DNA in *C. parvifithorax*, *D. melanogaster*, and *D. virilis*, respectively (Table 5). In summary, follicle cell underreplication and 16C/2C ratios are good predictors of satellite sequence and possibly heterochromatin content. Moreover, a clear trend exists in which larger genomes tend to have more underreplicated satellite DNA (Figure 5 and Table 5). For example, *D. virilis*, with the largest genome (364–438 Mb), has among the highest (40–48%) underreplicated content.

Conversely, *D. mojavensis*, with one of the smallest genomes (130–166 Mb), also has the least amount of underreplicated DNA (2–16%). In addition, some species, such as *D. pseudoobscura*, *D. melanogaster*, and *D. mojavensis*, exhibited a large range of intraspecific differences in underreplication (Table 5).

DISCUSSION

We provide the first systematic and replicated estimates of genome size and satellite DNA content in multiple species of *Drosophilidae*, revealing both intra- and interspecific differences in genome size. Of particular interest are the sequenced genomes of the 12 *Drosophila* species and how whole-genome sequence and accurate size estimates now allow us to more completely understand how these genomes have evolved and function.

Ploidy regulation and underreplication during oogenesis is conserved: Ploidy regulation in endoreduplicating ovarian follicle cells is evolutionarily conserved as all species we examined complete follicle cell DNA replication with 16C ploidy. Strict ploidy control appears critical for proper development of this tissue type. In *D. melanogaster*, hypomorphic mutations in the Rbf/E2F pathway allow ectopic DNA replication in follicle cells and disrupt ploidy control, but these mutations also lead to female sterility, indicating a more central function for Rbf/E2F than just control of ploidy (ROYZMAN *et al.* 1999; BOSCO *et al.* 2001; CAYIRLIOGLU *et al.* 2001). The evolutionary conservation of 16C follicle cell ploidy in all 38 species argues for a critical role for ploidy level in proper follicle cell function.

We also determined that all 38 species, and not just *D. melanogaster*, underreplicate their genomes in polyploid

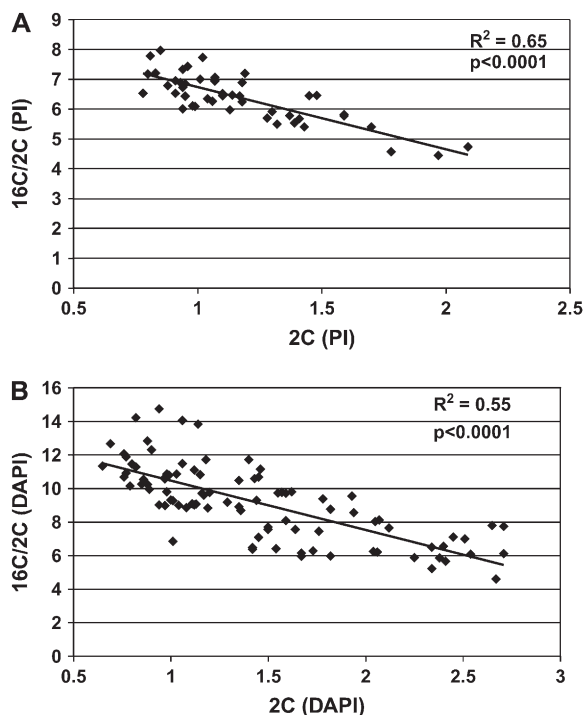


FIGURE 3.—The 16C/2C ratios are inversely proportional to 2C values. 16C/2C ratios were compared to their corresponding 2C values for PI (A) and DAPI (B) values. In each case, 16C/2C values decrease as genomes increase in size, indicating that a larger fraction of the genome is being underreplicated. A two-tailed P -value was calculated from the correlation coefficient (R) using Graphpad software. PI values (A) had 45 d.f. and DAPI values (B) had 90 d.f.

follicle cells. Underreplication also constitutes a conserved feature of all species examined in this study. Underreplication is a pervasive but poorly understood process with important implications for DNA replication fork barriers and transcription in diploid cells (LEACH *et al.* 2000; BELYAKIN *et al.* 2005). Structural features of heterochromatin satellite repeats, as opposed to specific sequences, have been proposed to act as replication barriers (LEACH *et al.* 2000). The fact that underreplication is conserved, despite great differences in satellite DNA content and species-specific repeat sequence motifs, implies that structural and possibly epigenetic factors act as fork barriers (DEMAKOVA *et al.* 2007). The availability of additional *Drosophila* genome sequences will allow a more thorough analysis of underreplicated genomic regions and genetic elements such as fork barriers that may control this conserved process.

Furthermore, by exploiting underreplication of satellite repeats, we detected surprising variation in satellite DNA content and in its contribution to genome size differences. The amount of underreplication fits well with cytological assays of heterochromatic regions as well as with previously described heterochromatin content estimates. Thus we propose that follicle cell underreplication values may be good predictors of heterochromatin content.

The variation in satellite DNA content and its significance for changes in genome size is consistent with previous ideas that genomes have expanded/contracted mainly by addition/deletion of repeat sequences (for review see HARTL 2000). In light of the 12 *Drosophila* genome sequences and our satellite DNA estimates, we can speculate as to the mechanisms by which these species have modified their satellite repeats. Unequal sister-chromatid exchange and replication errors have been suggested as possible molecular mechanisms that can produce variation in satellite DNA content (BRITTEN and KOHNE 1968; SOUTHERN 1975; SMITH 1976; STEPHAN and CHO 1994; PETROV 2001). However, unequal exchange of meiotic sister chromatids as well as replication errors are expected to give rise to both deletions and/or duplications. Unless meiotic drive or other species-specific selection acts upon these meiotic events, gametes bearing either deletions or duplications should be recovered in equal proportions, generating large intra- and interspecific variation. Our data suggest exactly the opposite: Intraspecific satellite DNA content differences are small whereas interspecific differences can be large (Table 5).

This raises an important question: Is genome size, and more specifically satellite DNA content, under selection? One obvious constraint on the contraction of heterochromatin repeats is centromeric function. In *D. melanogaster*, the minimum satellite DNA for a fully functional centromere has been measured to be ~ 420 kb (SUN *et al.* 1997). Other species are likely to have similar lower-limit constraints to ensure proper chromosome segregation. Among the species with the smallest genomes, *D. erecta*, *D. hydei*, *D. mercatorum*, and *D. mojavensis*, ~ 150 Mb or smaller (Table 3), none have $< 2\%$ satellite DNA (Table 5 and Figure 5A). Of the 12 sequenced species, we estimate *D. mojavensis* (strain 15081-1352.22) to have the smallest genome at 130 Mb and the least satellite DNA (2%). Interestingly, if the 2% satellite DNA (2.6 Mb) were distributed evenly among the six *D. mojavensis* chromosomes, then each chromosome would have ~ 430 kb of satellite heterochromatin. It will be informative to determine the chromosomal distributions of these repeats in different species and to ascertain whether these species adhere to the ~ 420 -kb limit seen in *D. melanogaster*.

Do *Drosophila* satellite and heterochromatin contents have upper limits? Several transcription factors have been shown to also bind satellite repeat sequences. Species-specific upper limits to heterochromatin content may be determined by threshold levels of euchromatic DNA-binding proteins that also bind satellite repeats (for review see ASHBURNER *et al.* 2005, p. 67). This model is attractive because it suggests that the species-specific genomic arrangements that place specific genes within the influence of heterochromatin dictates how much expansion/deletion is tolerated.

Estimates of total genome size and A:T content: Genome sizes for a number (20) of the *Drosophila* species examined here were reported upon in the earlier studies

TABLE 5
% of underreplication of heterochromatin satellite DNA

Species	PI		DAPI	
	Mean \pm SE (<i>n</i>)	Range	Mean \pm SE (<i>n</i>)	Range
<i>C. pararufithorax</i>	18 (1)	—	31 (1)	—
<i>C. procnemis</i>	30 (1)	—	30 (1)	—
<i>C. rufithorax</i>	21 (1)	—	31 (1)	—
<i>D. acutilabella</i>			22 \pm 0 (2)	22
<i>D. americana</i>	28 (1)	—	26 \pm 1 (2)	25–28
<i>D. ananassae</i>	21 \pm 2 (3)	17–23	25 \pm 0 (3)	25
<i>D. buskii</i>			22 (1)	—
<i>D. equinoxialis</i>			28 \pm 0 (2)	28
<i>D. erecta</i>	11 \pm 2 (2)	9–13	22 \pm 1 (2)	20–21
<i>D. funebris</i>			30 (1)	—
<i>D. grimshawi</i>	32 \pm 0 (1)	—	32 (1)	—
<i>D. guttifera</i>	3 \pm 2 (2)	1–5	19 \pm 2 (2)	17–21
<i>D. hydei</i>	1 (1)	—	22 \pm 2 (2)	21–24
<i>D. immigrans</i>	30 \pm 3 (2)	27–33	38 \pm 1 (3)	37–39
<i>D. littoralis</i>			26 (1)	—
<i>D. melanogaster</i>	26 \pm 3 (4)	19–33	28 \pm 1 (10)	23–40
<i>D. mercatorum</i>	12 (1)	—	19 \pm 3 (2)	16–22
<i>D. mimica</i>	27 \pm 1 (4)	25–30	35 \pm 1 (3)	34–37
<i>D. mojavensis</i>	8 \pm 4 (3)	2–16	20 \pm 1 (3)	17–22
<i>D. nanoptera</i>			37 \pm 2 (3)	34–41
<i>D. novamexicana</i>			26 \pm 1 (2)	25–27
<i>D. persimilis</i>	13 \pm 1 (3)	11–14	22 \pm 1 (3)	20–23
<i>D. pseudoobscura</i>	12 \pm 4 (3)	4–16	20 \pm 1 (3)	19–21
<i>D. repleta</i>			19 \pm 1 (3)	18–20
<i>D. sechellia</i>	24 \pm 0 (2)	23–25	27 \pm 0 (2)	26–27
<i>D. simulans</i>	20 \pm 1 (6)	14–23	28 \pm 3 (7)	23–38
<i>D. virilis</i>	44 \pm 1 (4)	40–48	42 \pm 1 (5)	40–46
<i>D. willistoni</i>	14 \pm 1 (3)	12–15	23 \pm 0 (3)	22–23
<i>D. yakuba</i>	21 \pm 2 (2)	19–23	32 \pm 9 (2)	23–41
<i>H. duncani</i>			30 \pm 0 (1)	—
<i>S. latifasciaeformis</i>			32 \pm 0 (2)	32
<i>S. lebanonensis</i>			25 \pm 0 (2)	25
<i>S. palmae</i>			25 \pm 0 (1)	—
<i>S. stonei</i>			26 \pm 0 (2)	25–26
<i>Z. badyi</i>			38 (1)	—
<i>Z. ghesquerei</i>			24 (1)	—
<i>Z. sepsoides</i>			46 \pm 5 (2)	39–53
<i>Z. tuberculatus</i>			35 \pm 2 (3)	31–43

Mean percentage of underreplication of satellite DNA in 16C follicle cells is shown as measured by PI and DAPI. Standard error (SE), the range (lowest and highest values), and the number of strains for each species (*n*) are shown.

mentioned above (Table 3). In many of those species, the genome sizes appear similar, although some deviate substantially. Unfortunately, the *Drosophila* species for which there were earlier genome size estimates came either from unpublished citations or from different investigations that used a wide range of methodologies or tissue types. Moreover, there is no strain origin information available for these estimates. Thus, for those species, the difference between previous estimates and ours cannot be evaluated (Table 3). For 12 of these species, our estimates differ by <50 Mb of previous values; we found two species (*D. americana* and *D. mojavensis*) to be ~50 Mb lower and four species (*D. buskii*, *D. equinoxialis*,

D. funebris, and *Scaptodrosophila lebanonensis*) to be ~50 Mb greater than previous estimates; we found two species (*S. latifasciaeformis* and *S. stonei*) to be ~90 Mb greater than previous estimates (Table 3). DAPI fluorescence alone was used in all six cases where our values are greater than previous reported estimates, and thus these higher values may not be as accurate as those previously determined by PI flow cytometry. We would predict therefore that these genomes are likely to have A:T-rich genomes because the DAPI values are higher than expected (Tables 3 and 5).

The smallest genome, 128 Mb for *D. mercatorum*, and the largest, 404 Mb for *D. virilis*, differ by as much as 3.2-fold

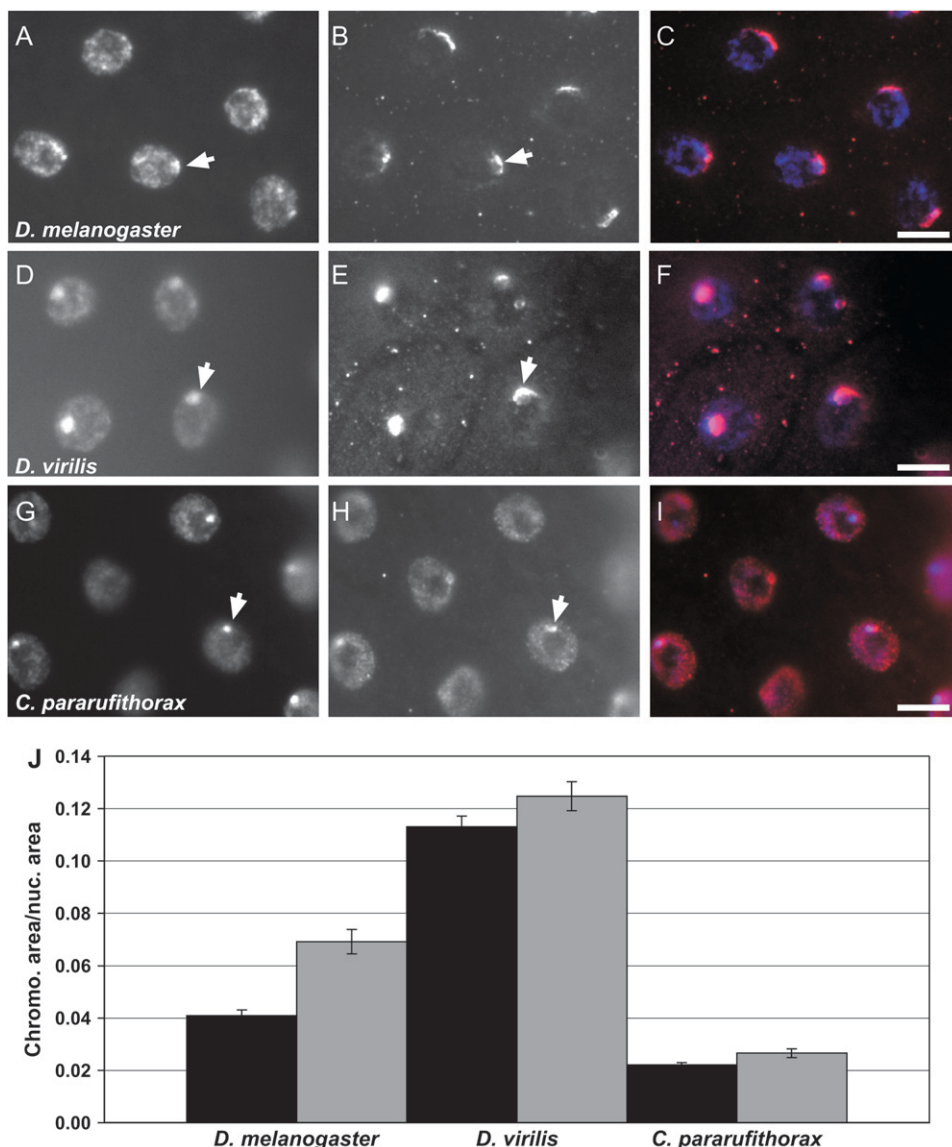


FIGURE 4.—Chromocenter size reflects satellite content. Stage 13 follicle cell nuclei were stained with DAPI (A, D, and G) and with antimethyl histone H3 (B, E, and H). Chromocenters (arrows) stain as a bright DAPI dot within the nucleus and are enriched for lysine-9 dimethyl H3. The merged images (C, F, and I) show colocalization of DAPI and lysine-9 dimethyl H3. Bars, 10 μ m. The area from each chromocenter was measured and normalized for nuclear area (J). Both DAPI (solid bars) and lysine-9 dimethyl H3 (shaded bars) area measurements show that, compared to *D. melanogaster*, *D. virilis* chromocenters are significantly larger ($P < 0.0001$) whereas *C. pararufithorax* has significantly smaller chromocenters ($P < 0.0001$). Standard error bars are shown and values represent averages from 35 cells (see MATERIALS AND METHODS).

(Table 3). Although our estimates suggest that up to 48% of *D. virilis* could be heterochromatic satellite DNA, this still does not account for the 3.2-fold difference in genome size with *D. mercatorum*. This difference is consistent, however, with a previous report that the *D. virilis* euchromatic genome has also expanded (MORIYAMA *et al.* 1998). By contrast, *D. virilis* is 1.6- to 1.9-fold larger than the 231-Mb genome of *D. grimshawi*, a difference that can be accounted for by an \sim 1.6-fold difference in satellite DNA estimates (Tables 3 and 5). In the close relatives *D. melanogaster*, *D. simulans*, and *D. erecta*, satellite DNA content differences are sufficient to explain the small but significant differences in our genome size measurements (Tables 3 and 5).

The importance of dye type is underscored by the genome size estimates for the Chymomyza species with PI *vs.* DAPI. While DAPI values are intrinsically less accurate when estimating total DNA content, they are, nevertheless, informative. It is noteworthy that some species

with relatively high 2C DAPI values, such as the three Chymomyza, did not have correspondingly high PI 2C values relative to *D. m. yw* (Table 3, supplemental Tables 1 and 2 at <http://www.genetics.org/supplemental/>). This suggests that the Chymomyza lineage is characterized by relatively high A:T-rich sequences. High Chymomyza A:T content could reflect high levels of AT-rich centric heterochromatin or indicate that Chymomyza euchromatin is more A:T rich than that of *D. melanogaster*. Chymomyza PI 16C/2C ratios (supplemental Table 1 at <http://www.genetics.org/supplemental/>) show no significant difference from the 16C/2C ratio observed for *D. m. yw*. Levels of underreplicating DNA in Chymomyza therefore appear similar to that of *D. melanogaster* (Table 5), and thus the relatively high A:T content in Chymomyza is likely a function of euchromatic, as opposed to heterochromatic, A:T sequences. Furthermore, cytological staining of at least one Chymomyza species chromocenter (Figure 4) indicates that high DAPI values are

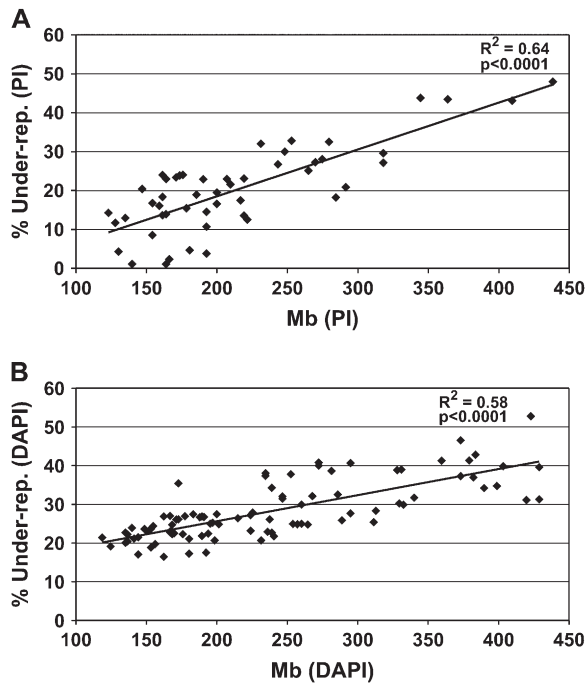


FIGURE 5.—Larger genomes have greater underreplication. The percentage of underreplication (y-axis) was calculated on the basis of 16C/2C values (see MATERIALS AND METHODS) and is shown plotted against haploid genome size in megabases, as determined by PI (A) and DAPI (B) flow cytometry. A trend line was added to show that, as genomes become larger, a greater fraction of the total DNA content is underreplicated. Note that the same trend is observed regardless of the dye used. Two-tailed *P*-values were calculated as in Figure 3.

likely due to A:T-rich euchromatin. Digestion with restriction enzymes that recognize either A:T- or G:C-rich sequences have confirmed that *Chymomyza* genomic DNA is more A:T rich than *D. melanogaster* and *D. virilis* (data not shown; P. CAMPBELL and G. BOSCO, unpublished data).

Species with both small and large genomes had DAPI/PI ratios >1, although the general trend was that larger genomes were more A:T rich (Figure 2B). This pattern is most apparent when comparing the relative 2C values derived from DAPI and PI in one of the largest genomes, *D. virilis* (Table 1, Figure 2B, supplemental Tables 1 and 2 at <http://www.genetics.org/supplemental/>). Because heterochromatin repeats are generally more A:T rich than euchromatic sequences (GALL *et al.* 1971), we conclude that material contributing to relatively high DAPI values is largely A:T-rich heterochromatin, except in the case of the *Chymomyza* discussed above.

A:T content and dye effects on small genomes: Genomes with DAPI/PI ratios <1, or relatively G:C-rich genomes, were all < ~200 Mb/haploid genome (Figure 2B). *D. persimilis*, *D. pseudoobscura*, and *D. simulans* were the most notable examples (supplemental Table 4 at <http://www.genetics.org/supplemental/>). These genomes exhibited significant underreplication, indicating that, although small, they still contain considerable amounts of satellite repeats. The question then arises as to the na-

ture of these repeats and why the DAPI/PI ratio is low. One possibility is that these genomes have low repeat content and repeats are not A:T rich. This is a likely explanation because DAPI 16C/2C ratios greater than the expected value of 8 were observed for smaller genomes (Figure 3B). DAPI thus appears to underestimate underreplication in smaller genomes because underreplicated satellite sequences are more G:C rich than satellite sequences of larger genomes. For example, in *D. melanogaster* >50% of satellite sequences (AAGAC, AAGAG, AAGAGAG, 1.688 satellite) are 28–40% G:C rich whereas the entire satellite sequences of *D. virilis* (ACAAACT, ATAAACT, and ACAAATT) are ~28% G:C rich (GALL *et al.* 1971; SCHWEBER 1974; LOHE and BRUTLAG 1986, 1987). For closely related sibling species, such as *D. melanogaster*, *D. simulans*, and *D. erecta* with nearly identical satellite repeat sequences, this trend is not apparent, although *D. melanogaster* contains more satellite repeats than *D. simulans*, which has more than *D. erecta* (Table 5), as previously described (LOHE and BRUTLAG 1987). Unfortunately, since the actual sequence identity of satellite repeats and their abundance in most species are unknown, a more thorough and inclusive analysis cannot be performed.

In one case, a *D. melanogaster* strain (BI 2057), we observed a discrepancy between the PI and the DAPI 2C values (Table 1). This suggested that this strain, unlike the other *D. melanogaster* strains, had acquired some additional A:T-rich DNA. However, the PI 16C/2C ratio (6.15) for this strain does not differ significantly from the other strains (Table 1). If additional A:T sequences are present, they are unlikely to consist of underreplicating satellite repeats. Without further molecular analysis it is difficult to say what might underlie the cause of this discrepancy.

We also found statistically significant differences in genome size among strains of a given species, although these differences in many cases were small. Examination of more strains from these species, especially strains freshly derived from nature, may be necessary to reveal more substantial differences. Specific examples with significant intraspecific variation in heterochromatin content have been described previously (HALFER 1981). Any phylogenetic analyses of genome size (G. BOSCO, T. MARKOW and B. McALLISTER, unpublished results) therefore will need to account for intraspecific variation as well as for the influence of dye.

The 12 *Drosophila* species genomes: Genomes of 12 *Drosophila* species have been sequenced, allowing us to compare the sizes of the euchromatic assembled portions of the sequenced genomes to sizes estimated with our methods and the contributions of heterochromatin to those sizes (Table 4). In four species, *D. ananassae*, *D. erecta*, *D. willistoni*, and *D. mojavensis* assembled genome sizes (<http://rana.lbl.gov/drosophila/> and DROSOPHILA 12 GENOMES CONSORTIUM 2007) are larger than those measured by flow cytometry. Size differences, when they

exist, are expected to be in the opposite direction: heterochromatin and satellite sequences should not be represented in the sequenced genomes and thus sequenced genomes should be the same or smaller than the estimates reported here. The largest discrepancy is in *D. mojavensis*, which has the lowest amount of underreplicated satellite DNA (Table 4). For *D. ananassae*, previous genome size estimates (ASHBURNER *et al.* 2005) are identical to ours, and our estimates do not differ with dye type, making it unlikely that this discrepancy reflects errors intrinsic to cytometric measurements of DNA content. In the case of these four species, it is possible that assembly sizes do not accurately represent euchromatic genome sizes as assembly errors have been reported for previous genome releases, including *Drosophila*, mouse, and human genomes (BENOS *et al.* 2001; CELNIKER *et al.* 2002; CHEUNG *et al.* 2003a,b).

The differences in genome size and heterochromatin content point to specific and testable evolutionary questions. For example, is loss and or gain of heterochromatic repeat elements the same for different repeat types and for different chromosomes, as has been shown for *D. melanogaster* and closely related species? Surprisingly little is known about the repeat sequences, abundance, and distribution of satellite sequences in all but a handful of *Drosophila* species. What are the costs, if any, of the possession of higher amounts of heterochromatin in one *vs.* another strain of the same species? In *D. melanogaster*, varying amounts of heterochromatin such as Y chromosome translocations have been shown to be a potent suppressor of position-effect variegation, thus raising the question as to how different strains and different species with vast differences in heterochromatin could use or cope with large differences (BECKER 1977). Aside from the known structural roles that heterochromatin plays in centromere function (SUN *et al.* 1997, 2003) and meiotic chromosome pairing (HAWLEY *et al.* 1993; DERNBURG *et al.* 1996), are there other important functions for heterochromatin, such as epigenetic modification, that are under selection and possibly driving genome expansion? Our genome size and heterochromatin estimates complement the *Drosophila* genome sequences and will allow a more in-depth exploration of the possible mechanisms and evolutionary forces by which genomes have expanded and contracted.

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