

The *Drosophila* BAC resource

1 **The 19 Genomes of *Drosophila*: a BAC Library Resource for Genus-wide and**
2 **Genome Scale Comparative Evolutionary Research**

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ABSTRACT

40 The genus *Drosophila* has been the subject of intense comparative phylogenomics
41 characterization to provide insights into genome evolution under diverse biological and
42 ecological contexts, and to functionally annotate the *D. melanogaster* genome, a model
43 system for animal and insect genetics. Recent sequencing of 11 additional *Drosophila*
44 species from various divergence points of the genus is a first step in this direction.
45 However, to fully reap the benefits of this resource, the *Drosophila* community is faced
46 with two critical needs: i.e. the expansion of genomic resources from a much broader
47 range of phylogenetic diversity, and the development of additional resources to aid in
48 finishing the existing draft genomes. To address these needs, we report the first synthesis
49 of a comprehensive set of BAC resources for 19 *Drosophila species* from all three
50 subgenera. Ten libraries were derived from the exact source used to generate 10 of the 12
51 draft genomes, while the rest were generated from a strategically selected set of species
52 based on salient ecological and life history features, and their phylogenetic positions. The
53 majority of the new species have at least one sequenced reference genome for immediate
54 comparative benefit. This 19 BAC library set was rigorously characterized and shown to
55 have: large insert sizes (125 to 168 kb), low non-recombinant clone content (0.3% to
56 5.3%), and deep coverage (9.1X - 42.9X). Further, we demonstrated the utility of this
57 BAC resource for generating physical maps of targeted loci, refining draft sequence
58 assemblies, and identifying potential genomic rearrangements across the phylogeny.

59

60

INTRODUCTION

61 The genus *Drosophila* contains approximately 2000 species of diverse morphology,
62 ecology and behavior that are placed in three major lineages: subgenus *Sophophora*,
63 subgenus *Drosophila* and subgenus *Dorsilopha* (Markow and O'Grady 2006, 2007). The
64 most widely studied species in the genus, *D. melanogaster*, is firmly established as the
65 premier model system for many biological research areas such as neurobiology, medicine
66 and population biology (Rubin and Lewis 2000). Several other species in this genus,
67 such as *D. pseudoobscura* and *D. virilis*, have also been utilized as genetic model systems
68 particularly for evolutionary studies (Anderson *et al.* 1991; Popadic and Anderson 1994;
69 Orr and Coyne 1989; Charlesworth *et al.* 1997; Vieira *et al.* 1997; Sweigart 2010).
70 Recently, the genomes of *D. melanogaster* and 11 other *Drosophila* species, whose most
71 recent common ancestor occurred more than 45-50 million years ago, have been
72 sequenced, assembled and annotated (Adams *et al.* 2000; Myers *et al.* 2000; Celniker *et*
73 *al.* 2002; Richards *et al.* 2005; *Drosophila* 12 Genomes Consortium 2007; Gilbert 2007).
74 Species were selected for genome sequencing partly based on their relationship with *D.*
75 *melanogaster*. Nine of the twelve sequenced genomes were sampled from one subgenus,
76 *Sophophora*, to which *D. melanogaster* belongs and the remaining three are from the
77 *Drosophila* subgenus. These sequences have already greatly improved understanding of
78 the evolution and regulation of eukaryotic genes and genomes through comparative
79 analyses (Stark *et al.* 2007). However, to fully reap the benefits from this unique resource,
80 the *Drosophila* community has faced with two critical needs: first, the development of
81 additional genomics resources to aid in finishing the 11 existing draft genome sequences;

82 and second, the generation of additional genomic resources that encompass a much
83 broader range of phylogenetic diversity.

84 Towards this direction, we constructed a comprehensive set of bacterial artificial
85 chromosome (BAC) libraries for 19 different *Drosophila* species representing a broad
86 spectrum of phylogenetic diversity. BAC libraries are powerful tools for comparative
87 genome research (Kim *et al.* 1996; The International Human Genome Mapping
88 Consortium 2000a, b; Hoskins *et al.* 2000; Locke *et al.* 2000; Osoegawa *et al.* 2000, 2001,
89 2004; Gregory *et al.* 2002; Eichler and DeJong 2002; Gibbs *et al.* 2003; Krzywinski *et al.*
90 2004; Gonzalez *et al.* 2005; Ammiraju *et al.* 2006; *Drosophila* 12 Genomes Consortium
91 2007; Kim *et al.* 2008; Murakami *et al.* 2008) especially in taxa containing highly
92 repetitive genomes (Ellison and Shaw 2010; Havlak *et al.* 2004; Fang *et al.* 2010).
93 Genome sequences are available for 10 of 19 species for which BAC libraries are
94 constructed, some of which were instrumental in facilitating sequence assemblies
95 (*Drosophila* 12 Genomes Consortium 2007), and they remain a high priority resource for
96 improving and finishing several of the low coverage draft genome assemblies. BAC
97 libraries for species without sequenced genomes present an important resource for
98 positional cloning and large-scale targeted comparative genome analyses.

99 We selected 19 species within three lineages of the genus *Drosophila* for BAC library
100 construction (Figure 1). These species shared a common ancestor approximately 40-60
101 million years ago (Powell 1997) and were selected because of their varied evolutionary
102 distances from *D. melanogaster* and other sequenced species, their diverse ecologies and
103 life history characters, and the fact that they can be reared in the laboratory and used in
104 experimental work in the future. Ten BAC libraries were constructed as a resource for

105 generating BAC end mate-pair sequence to assist in the assembly of whole-genome
106 shotgun sequences, and for enabling future genomic research (*Drosophila* 12 Genomes
107 Consortium 2007). Beyond those 10 species, we are interested in generating BAC library
108 resources for representative species of lineages not yet targeted for sequencing but which
109 fill in large phylogenetic gaps. The majority of these species have at least one previously
110 sequenced reference genome for immediate comparative benefit. In addition, this new set
111 of species facilitates the “ladder and constellation” approach of modified phylogenetic
112 shadowing proposed by Clark *et al.*
113 (http://flybase.org/static_pages/news/whitepapers/GenomesWP2003.pdf) for annotating
114 genome data. In this approach ladder rungs constitute successively increasing divergence
115 points and constellations are clusters of species attaching to these divergence points. This
116 set of 19 BAC libraries documented here will further advance the genus *Drosophila* as an
117 ideal eukaryotic comparative genomics system designed to: 1) provide sequencing
118 resources for comparative annotation of the *D. melanogaster* genome; and 2) provide
119 genomic resources for experimental investigation of gene function throughout the genus
120 *Drosophila*.

121

122 MATERIALS AND METHODS

123

124 Fly culturing and embryo collection

125 Fly cultures were expanded on banana/opuntia medium
126 (<http://flyfood.arl.arizona.edu/opuntia.php3>) and healthy sexually mature adult flies were
127 introduced into plexiglass oviposition chambers kept on a 16:8 light/dark cycle at 24-

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128 25°C with a relative humidity of 60-80%. Exceptions to this procedure were: *D. littoralis*,
129 *D. novamexicana*, *D. americana*, *D. grimshawi* and *D. persimilis* cultures which were
130 oviposited at 20-22°C, whereas *D. albomicans* was oviposited at 17°C. Medium for *D.*
131 *sechellia* was supplemented with 0.5% (v/v) hexanoic acid and 0.5% (v/v) octanoic acid
132 to stimulate oviposition. Oviposition medium for *D. grimshawi* was supplemented with
133 2% (w/v) methylparaben to prevent overgrowth of fungus. *Drosophila busckii* and *D.*
134 *grimshawi* cultures were grown on Wheeler-Clayton medium
135 (<http://flyfood.arl.arizona.edu/wheeler.php3>). *Drosophila grimshawi* adults were
136 separated by sex until day of placement in the oviposition chamber to enhance embryo
137 production. Adult flies were allowed to oviposit on a given plate for as long as possible
138 without larval hatch. This interval varied between four and 48 hours depending on the
139 species. About 1.2-1.5 grams wet weight embryos were pooled in batches and stored at -
140 80°C at the end of each oviposition session.

141

142 **Nuclei preparation and BAC library construction**

143 Embryos were gently homogenized in PBS buffer (0.76% NaCl, 4mM NaH₂PO₄,
144 9mM Na₂HPO₄, PH 7.0) using a Dounce Tissue Grinder (Wheaton Science), centrifuged
145 at 4 °C at 1,430g for 15 min and resuspended in PBS buffer. The suspension was then
146 mixed with an equal volume of 1% InCert Agarose (CAMBREX, in PBS buffer) at 45 °C
147 and transferred into plug molds. Treatment of plugs to produce un-sheared megabase-size
148 DNA was as described (Luo and Wing 2003). BAC libraries were constructed as
149 previously described (Luo and Wing 2003; Ammiraju *et al.* 2006).

150

151

152 **BAC library characterization**

153 DNA from a random sample of 260-480 BAC clones from each library was
154 isolated, restriction digested with *NotI*, and run on CHEF gels for insert size
155 determination as previously described (Luo and Wing 2003; Ammiraju *et al.* 2006).

156 High colony density hybridization filters for each library were prepared using
157 Genetix Q-bots (Genetix) as described previously (Luo *et al.* 2006; Ammiraju *et al.*
158 2006). Nine gene specific probes were chosen that represented all chromosomes of *D.*
159 *melanogaster* (Tables S1 and S2). All probe DNA fragments were PCR amplified from
160 the *D. mojavensis* genome and gel purified using a QIAEX II (Qiagen) kit. Table S1 lists
161 the primer sequences used for each probe. Purified DNA fragments were sequenced and
162 similarity searches were conducted to validate their specificity. Probes were prepared by
163 labeling with ³²P dCTP using a DecaprimeII random prime labeling kit (Ambion), and
164 hybridizations were carried out as described by Ammiraju *et al.* (2006). Positive clones
165 were picked, re-arrayed on to colony filters, followed by a secondary hybridization with
166 individual probes.

167

168 **Fingerprinting and contig assembly**

169 Positive hybridization clones were fingerprinted using SNaPshot (Luo *et al.* 2003;
170 Kim *et al.* 2008), and assembled into contigs with FPC v 8.5.2 (Soderlund *et al.* 2000;
171 www.agcol.arizona.edu) at a fixed tolerance value 4 and an initial Sulston score $1e^{-50}$
172 (Ammiraju *et al.* 2006)

173

174 **BAC end sequencing and in silico analysis**

175 Fingerprinted BAC clones were end sequenced with a universal T7 primer (5' TAA
176 TAC GAC TCA CTA TAG GG 3') and a custom primer BES_HR (5' CAC TCA TTA
177 GGC ACC CCA 3') following previously described methods (Kim *et al.* 2008). BAC end
178 sequences (BES) were submitted to GenBank with the following accession numbers: *D.*
179 *simulans* (EI211963.1-EI212067.1), *D. sechellia* (CZ549016.1-CZ549204.1), *D. yakuba*
180 (EI89369.1-EI189559.1), *D. erecta* (CZ548656.1-CZ548834.1), *D. ananasseae*
181 (CZ548467.1- CZ548655.1), *D. persimilis* (EI188778.1-EI189177.1), *D. willistoni*
182 (EI189178.1- EI189368.1), *D. americana* (EI189178.1-EI189368.1), *D. novamexicana*
183 (DU169152.1-DU169329.1), *D. virilis* (CZ549205.1-CZ549371.1), *D. littoralis*
184 (EI211597.1-EI211779.1), *D. repleta* (EI211780.1-EI211962.1), *D. mercatorum*
185 (EI188452.1-EI188610.1), *D. mojavensis* (CZ548835.1-CZ549015.1), *D. arizonae*
186 (EI211417.1-EI211231.1), *D. hydei* (EI188451.1-EI188450.1), *D. grimshawi*
187 (EI188111.1-EI188299.1), *D. albomicans* (EI211043-EI211230.1), and *D. busckii*
188 (EI211418.1-EI211596.1).

189 All BESs were masked with Repeat Masker (version3.1.0) against a redundant repeat
190 database with sequences obtained from fly base (www.FlyBase.org) and Repbase
191 (www.girinst.org). These sequences were used to conduct BLAST analysis against the
192 mitochondrial (NC_001709, 19517 bp) and nuclear genome sequences of *Drosophila*
193 *melanogaster* (Build 5.1) and the freeze 1 genome assemblies from the remaining eleven
194 species (<http://rana.lbl.gov/Drosophila/caf1.html>, and
195 <http://insects.eugenes.org/species/data/>). To compensate for the lack of whole genome
196 sequences and to minimize the bias of sequence divergence, the genome sequences of *D.*

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197 *virilis* and *D. mojavensis* were used as pseudo-reference sequences for the *D. virilis* and
198 *D. repleta* species group, respectively. BES from *D. albomicans* and *D. busckii* was
199 compared to the *D. grimshawi* sequences.

200 In addition, similarity searches were conducted with complete gene sequences of each
201 probe against the 12 *Drosophila* whole genome sequences (*Drosophila* 12 Genomes
202 Consortium 2007). Homologs with a minimum alignment length of 100 bp and 75 % of
203 nucleotide identity were retained for further analysis and for a comparison of their
204 presence or absence in FPC derived contigs.

205

206

RESULTS AND DISCUSSION

207

***Drosophila* strain selection and genome sizes:**

209 Several criteria were used for careful evaluation of the different *Drosophila*
210 species strains used for BAC resource development in this study. First, all fly lines were
211 inbred for a minimum of 8 generations by sib-sib mating to reduce the extent of
212 heterozygosity and subsequently sequenced at six nuclear loci to verify homozygosity
213 (data unpublished). Second, to minimize endosymbiont contamination (*Wolbachia* spp.
214 and *Spiroplasma* spp.) at least 5 adult fly DNA samples from each species were screened
215 with established protocols (Mateos *et al.* 2006). Finally, species identity was confirmed
216 by both morphological and molecular approaches. When a suitable nuclear or
217 mitochondrial DNA marker was known for a species, that marker was amplified,
218 sequenced and validated. Additionally, salivary gland chromosomes from third instar
219 larvae were prepared and inspected for inversion polymorphism microscopically. Only

220 homokaryotypic lines were used. All strains (Table 1) are deposited in the UC San Diego
221 *Drosophila* Stock Center and are publicly available as a community resource.

222 Genome size of an organism is the most important factor in determining the depth
223 of a genomic library (reviewed in Gregory 2005). Previously determined genome sizes
224 (Bosco *et al.* 2007) were used in this study for estimating the coverage of the BAC
225 libraries for different *Drosophila* species. Bosco *et al.* (2007) employed two nucleic-acid
226 binding fluorescent dyes, propidium iodide (PI) and 4',6-diamidino-2-phenylindole
227 (DAPI), in conjunction with flow cytometry to determine genome sizes of 38 species of
228 Drosophilidae, including the 12 sequenced *Drosophila* species (*Drosophila* 12 Genomes
229 Consortium 2007).

230 The genome sizes of 15 of the 19 *Drosophila* species used in this study were
231 based on the PI method and the remaining species (*D. novamexicana*, *D. littoralis*, *D.*
232 *repleta* and *D. busckii*) genome sizes were based alone the DAPI method alone (for
233 which the PI data was not available) (Table 1). Nine of the *Drosophila* species strains
234 were not the same as the strains analyzed by Bosco *et al.* (2007). An important finding to
235 consider, as reported by Bosco *et al.* (2007) and Gregory and Johnston (2008), is that
236 DAPI may overestimate genome size which could affect the estimated genome coverage
237 of these 4 libraries.

238 Genome sizes of two species, *D. arizonae* and *D. albomicans*, were not known, so
239 the genome sizes of closest relatives *D. mojavensis* and *D. immigrans*, respectively were
240 applied to estimate the tentative genome coverages of their respective BAC libraries. The
241 genome sizes among the 19 *Drosophila* species varried by ~3.2 fold, with the smallest
242 being *D. mercatorum* and the largest *D. virilis* (Table 1).

243

244 **BAC library construction and characterization:**

245 Three different restriction enzymes were used for BAC library construction:
246 *HindIII*, *BamHI*, and *BstYI*. Fifteen of the 19 libraries were constructed from DNA
247 partially digested with *HindIII*, followed by size selection and ligation into the *HindIII*
248 site of pIndigoBAC536*SwaI* (Ammiraju *et al.* 2006) (Table 1). Two libraries each were
249 generated similarly from *BamHI* (*D. ananassae* and *D. mojavensis*) and *BstYI* (*D. virilis*
250 and *D. americana*) restriction digests. All libraries, except for the *D. busckii* library (two
251 ligations) were built from single ligations. The number of clones in the 19 BAC library
252 set ranged between 11,520 to 55,296 (Table 1), which were arrayed into 384-well
253 microtiter plates for long-term storage in -80°C freezers at the Arizona Genomics
254 Institute's (AGI) BAC/EST Resource Center (www.genome.arizona.edu).

255 Insert sizes of individual clones in each library ranged from 10 kb to 371 kb, with
256 the majority over 120 kb (Figure 2). The average insert sizes of these libraries ranged
257 from 125 to 168 kb (Table 1). Percentages of non-insert containing clones ranged
258 between 0.3% - 5.3%, which is typical for BAC libraries constructed at AGI (Ammiraju
259 *et al.* 2006).

260

261 **Genomic redundancy of the *Drosophila* BAC libraries**

262 We estimated the genomic depth of the 19 *Drosophila* BAC set by three different, but
263 complementary approaches. First, we estimated the redundancy of each library
264 empirically from the average insert size, total number of clones, and the genome size of
265 the corresponding lineage, which ranged approximately between 5.7 - 32.8 fold (Table 1).

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266 To assess the randomness and extent of representational heterogeneity for different
267 genomic regions, we screened the entire set of 19 *Drosophila* BAC libraries with 9 gene
268 specific probes in two successive rounds of hybridizations (methods; Tables S1 and S2).

269 In brief, 4196 putative positive BAC clones were identified in the first round of
270 hybridization, 3809 (91%) were confirmed by a second hybridization. The number of
271 positive hits per library ranged from 1 to 108 (Table S3). At least one positive hit per
272 each probe was detected for all the libraries with the exceptions of the *D. americana*, *D.*
273 *repleta*, *D. hydei* libraries for probe X-CG11387 and *D. ananassae* for probe 3R-
274 CG31247 (Table S3). In these four species no hits were found, even upon three rounds of
275 library screening, with different hybridization stringencies. For *D. ananassae*, the whole
276 genome draft sequence was available (<http://rana.lbl.gov/Drosophila/caf1.html>), and
277 similarity searches revealed the presence of the probe sequence (3R-CG31247; Table S2)
278 in the draft sequence assembly. Therefore, at least in the case of *D. ananassae*, it appears
279 that methodological and/or library coverage issues prevented recovery of this gene via the
280 hybridization based approach, possibly due to use of heterologous probes, multiple usage
281 of high density colony filters, or cloning bias (under and over representation of genomic
282 regions due to usage of a single restriction enzyme during library construction). More
283 data is required to confirm the absence of the gene X-CG11387 in other three species (*D.*
284 *americana*, *D. repleta*, *D. hydei*).

285 Hybridization based genome coverage's ranged from 9.1X (*D. americana*) to
286 42.9X (*D. hydei*). In only two species, *D. mercatorum* and *D. willistoni*, the hybridization
287 based coverage was slightly lower than expected (Table 2). The remaining 17 libraries
288 either had nearly equal or higher coverage than predicted (Table 2, Table S3). The *D.*

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289 *albomicans* BAC library showed a ~3.6 fold higher than expected coverage based on
290 hybridization (Table 2), which could have resulted from not having accurate genome size
291 estimation for this species (Table 1).

292 A third and a more rigorous approach using fingerprinted contig (FPC) based
293 estimations of genomic redundancy of BAC libraries was applied using a similar strategy
294 as our previous analysis of a set of 11 *Oryza* (cultivated and wild rice) BAC libraries
295 (Ammiraju *et al.* 2006). This approach can discriminate the unavoidable cloning bias
296 from those of cross hybridizations and genetic rearrangements such as duplications. All
297 3809 hybridization derived BAC clones were fingerprinted and 3005 (79%) successful
298 fingerprints were assembled into physical contigs (Tables S4 and S5). Under a scenario
299 of single copy probes and one contig per probe for each species, the theoretically
300 expected number of contigs is 171 (9 probes for 19 libraries). However several
301 exceptions were found; a) as described above, 1 probe X-CG11387 had no hits in the *D.*
302 *americana*; *D. repleta* and *D. hydei* libraries, and another probe - 3R-CG31247 - had no
303 hits in the *D. ananassae* library (Table S3); b) clones detected from 6 hybridizations (*D.*
304 *yakuba*, *D. persimilis* and *D. willistoni* with probe X-CG11387; *D. mercatorum* with
305 probe 2L-CG4128; *D. mercatorum*, *D. grimshawi* with probe 4-CG2999) resulted in the
306 presence of singletons (Table S5) (all these instances resulted in less than 3 positive
307 clones, Table S3). Taking into account the absence of these contigs in these species, 161
308 contigs are expected.

309 Our FPC analysis revealed a total of 211 contigs, 50 additional contigs than the
310 expected number of 161 (Table S4). The number of contigs and respective coverage
311 differed among different *Drosophila* libraries for the same probe (Table S5). Five probes

312 (X-CG11387, X-CG32611, 3L-CG10948, 3R-CG31247, 4-CG2999) essentially behaved
313 as single copy probes in most *Drosophila* libraries (Table S5). The remaining four
314 detected on average, 1.4 or more contigs/per probe (Table S5). To better understand if
315 these deviations from expectation (50 additional contigs) were due to technical issues
316 (cross hybridization and assembly artifacts,) and/or lineage specific genetic changes, we
317 gathered data from two additional experiments. First, based on BES mapping
318 information (methods), we classified 142 contigs as primary (those that map to the
319 expected genomic location) and 69 additional contigs as secondary (27 contigs that
320 cannot be positioned in any genome and 42 contigs that map to non-orthologous
321 locations), a good agreement between the results of FPC analyses and mapping
322 information (Tables S2 and S6).

323 Second, nucleotide and protein similarity searches of the probe (or gene) sequences
324 revealed that several secondary sites (17/42 secondary contigs) contained small cross
325 hybridizing paralogous sequences (Table S6, indicated with *). It is possible that the 25
326 remaining secondary sites also contained very small cross hybridizing sequences that
327 were not easily detected through similarity searches. In addition, sequence analysis of
328 the extended flanking sequences of the primary sites with the secondary sites revealed no
329 evidence of synteny, suggesting cross hybridization as the main cause for these additional
330 contigs.

331 To provide a conservative estimate of genome coverage, we considered each identified
332 contig as an independent locus and calculated a weighted FPC coverage that accounts for
333 the presence of several loci (Table S4; Ammiraju *et al.* 2006). Estimated FPC coverage

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334 for the 19 libraries (Table 2 and Table S4), ranged between 7 to 37X. Only two libraries
335 had coverage below nine fold: *D. willistoni* (7X) and *D. americana* (8X).

336 Twelve libraries showed a ratio close to 1:1 between the FPC and empirically
337 estimated coverage (Table 2). The *D. willistoni*, *D. littoralis*, *D. repleta*, *D. mercatorum*,
338 *D. mojavensis*, *D. arizonae* and *D. busckii* libraries showed ratios equal or below 0.7:1
339 (Table 2; Table S4). The difference between hybridization based and contig based
340 estimates of library coverage is due to the difference in the number of loci used to
341 calculate the coverage. While each probe is considered as a single locus in the
342 hybridization based approach, each secondary contig is considered as an independent
343 locus in the FPC based approach (Table 2; Tables S3 and S4). Together, these results
344 showcase the high quality and deep representational coverage of each of 19 *Drosophila*
345 genomes in their respective libraries.

346

347 **Utilization of BAC libraries**

348 Although a few *Drosophila* BAC libraries have already been reported in the literature
349 (Hoskins *et al.* 2000; Locke *et al.* 2000; Gonzalez *et al.* 2005; Osoegawa *et al.* 2007;
350 Murakami *et al.* 2008), this is the first synthesis and characterization of a comprehensive
351 set of BAC library resources for the genus, which fills a critical void for the *Drosophila*
352 research community. Hybridization of nine different probes to the full set of libraries
353 demonstrates the feasibility of isolating homologous regions across the entire genus.
354 Combined with high-throughput sequencing methods (Wicker *et al.* 2006), this set of
355 libraries provides an excellent resource for comparative studies of targeted genomic
356 regions (e.g., Leung *et al.* 2010).

357 First, BAC libraries from species that do not yet have a reference genome sequence
358 themselves provide a source for identifying genome rearrangements in comparisons with
359 the available genome sequences. For example, end sequences of BACs isolated with the
360 X-linked probe CG32611 from *D. novamexicana* map at an unexpected position within
361 contig 12970 of *D. virilis*, indicating a putative small inversion at the base of the X
362 chromosome that had not been previously identified (Vieira *et al.* 1997). Another putative
363 inversion was also revealed in *D. arizonae* by the localization of end sequences of clones
364 hybridizing to CG3139 in the genome sequence of *D. mojavensis*. Targeted analyses
365 inversion breakpoints are also enabled by the availability of these BAC libraries and
366 informed by the reference genome sequences. Evans *et al.* (2007) used cytological
367 evidence on the position of an inversion in *D. americana* to develop probes for isolating
368 its breakpoints from the respective BAC clones. In addition, the BAC libraries for the
369 nine un-sequenced *Drosophila* species provide robust templates for the whole genome
370 physical and sequence frameworks. In this direction, the entire *D. persimilis* BAC library
371 was fingerprinted, bidirectionally end sequenced, and assembled into a whole genome
372 physical map. This map was aligned to the *D. persimilis* and *D. pseudoobscura* draft
373 sequences, and is currently under editing (data not shown).

374 An extremely important application of the BAC resources reported here is in the
375 ability to use functional genomics to test genes underlying the differences between
376 *Drosophila* species. The tool kit for functional analyses of *Drosophila* has taken a major
377 leap forward with the recent establishment of the P/ΦC31 artificial chromosome
378 manipulation (P[acman]) transgenesis platform (Venken *et al.* 2006, 2007, 2009). While
379 still reliant on the P transposable element for transformation, this BAC transgenic system

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380 significantly improves upon the size of the DNA to be carried in the vector (>130 Kb),
381 and it's site specific integration in the fly genome. An important feature of the P[acman]
382 system is recombining – which permits cloning/transfer of large DNA fragments from
383 existing *Drosophila* P1 or BAC clones through a homologous recombination mediated
384 gap repair process. Therefore, a combination of the P[acman] system with the 19
385 *Drosophila* BAC libraries will provide an unprecedented opportunity to the fly
386 community to access, transfer and manipulate virtually any genomic region of interest
387 (large genes or even gene clusters) covering the entire phylogenomic range of the genus
388 *Drosophila*.

389 Finally, the BAC library set reported here can be used to further improve many of the
390 existing *Drosophila* draft sequence assemblies (*Drosophila* 12 Genomes Consortium
391 2007), and aid in the characterization of lineage specific rearrangements. For example,
392 physical mapping of BAC contigs, or individual BAC clones, identified by hybridization
393 probes designed from draft *Drosophila* genome sequences, has revealed and confirmed
394 chromosomal location of several sequence contigs from the draft assemblies, as well as
395 their relationship to *D. melanogaster* (Table S6). Conserved linkage and physical markers
396 were used to infer the physical organization of the assembled genome assemblies relative
397 to reference chromosome maps (Schaeffer *et al.* 2008), and these BAC libraries serve as
398 an appropriate resource to isolate regions at inferred gaps between adjacent contigs (e.g.,
399 Hoskins *et al.* 2000). Using hybridization to recover genome regions containing target
400 genes, combined with end sequencing of positive clones further reveals the conserved
401 linkage among *Drosophila* species. For example, scaffolds 20 and 24 map to X[A], 29 to
402 3L[D] and 30 to 4[F] in *D. sechellia*, 4512 4[F] in *D. erecta*, 12984 3R[B] and 12947

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403 4(LR)[F] in *D. ananassae*, 48 XR[D/A] and 103 5[F] in *D. persimilis*, 5 group M 5[F] in
404 *D. pseudoobscura*, 13052 6[F] in *D. virilis* (*Drosophila* 12 Genomes Consortium, 2007),
405 6498 6[F] in *D. mojavensis* and 14822 6[F] in *D. grimshawi* (Table S6).

406 These libraries are likely to facilitate a wide array of comparative, evolutionary and
407 functional genomics studies and play a major role in advancing the *Drosophila* biology.

408

409

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FIGURE 1. Phylogenetic tree of 19 species and *D. melanogaster* selected for the *Drosophila* BAC resource project. The phylogenetic relationships and approximate divergence times among the *Drosophila* species in our study were determined from a compilation of prior analyses (Pitnick *et al.* 1995; Markow and O'Grady 2006; *Drosophila* 12 Genomes Consortium 2007).

FIGURE 2. Insert size distribution of 19 *Drosophila* BAC libraries. Histograms A to S depict the insert size distribution in the 19 different libraries. For each histogram, X axis represents insert size (kb) and Y axis represents the number of clones in a particular insert size range.

A: *D. simulans* (DS_ABa); Average Insert Size 158 kb; B: *D. sechellia* (DS__Ba); Average Insert Size 139 kb; C: *D. yakuba* (DY__Ba); Average Insert Size 148 kb; D: *D. erecta* (DE_TBa); Average Insert Size 149 kb; E: *D. ananassae* (DA__Ba); Average Insert Size 148 kb; F: *D. persimilis* (DP__Ba); Average Insert Size 151 kb; G: *D. willistoni* (DW__Ba); Average Insert Size 150 kb; H: *D. americana* (DA_ABa); Average Insert Size 136 kb; I: *D. novamexicana* (DN__Ba); Average Insert Size 155 kb; J: *D. virilis* (DV_VBa); Average Insert Size 127 kb; K: *D. littoralis* (DL__Ba); Average Insert Size 168 kb; L: *D. repleta* (DR__Ba); Average Insert Size 143 kb; M: *D. mercatorum* (DM__Ba); Average Insert Size 125 kb; N: *D. mojavensis* (DM_CBa); Average Insert Size 143 kb; O: *D. arizonea* (DA_CBa); Average Insert Size 133 kb; P: *D. hydei* (DH__Ba); Average Insert Size 146 kb; Q: *D. grimshawi* (DG__Ba); Average Insert Size 127 kb; R: *D. albomicans* (DA_BBa); Average Insert Size 130 kb; S: *D. busckii* (DB__Ba); Average Insert Size 166 kb.

TABLE 1 Characteristics of the 19 *Drosophila* BAC library set

Species	Group ^e	Stock number ^f	Library name	Enzyme	Genome size (Mb)	Average insert size (Kb)	Clone number	Calculated genome coverage ^d
<i>D. simulans</i>	MEL	DSSC# 14021-0251.195	DS_ABa	<i>Hind</i> III	160 ^a	158	18432	18.2
<i>D. sechellia</i>	MEL	DSSC # 14021-0248.25	DS__Ba	<i>Hind</i> III	166 ^a	139	18432	15.4
<i>D. yakuba</i>	MEL	DSSC# 14021-0261.01	DY__Ba	<i>Hind</i> III	188 ^a	148	11520	9.1
<i>D. erecta</i>	MEL	DSSC #14021-0224.01	DE_TBa	<i>Hind</i> III	145 ^a	149	18432	18.9
<i>D. ananassae</i>	MEL	DSSC # 14024-0371.13	DA__Ba	<i>Bam</i> HI	215 ^a	148	36864	25.4
<i>D. persimilis</i>	OBS	DSSC# 14011-0111.49	DP__Ba	<i>Hind</i> III	183 ^a	151	18432	15.2
<i>D. willistoni</i>	WIL	DSSC# 14030-0811.24	DW__Ba	<i>Hind</i> III	206 ^a	150	18432	13.4
<i>D. americana</i>	VIR	DSSC #15010-0951.15	DA_ABa	<i>Bst</i> YI	275 ^a	136	11520	5.7
<i>D. novamexicana</i>	VIR	DSSC# 15010-1031.14	DN__Ba	<i>Hind</i> III	244 ^b	155	13440	8.5
<i>D. virilis</i>	VIR	DSSC # 15010-1051.87	DV_VBa	<i>Bst</i> YI	404 ^a	127	55296	17.4
<i>D. littoralis</i>	VIR	DSSC# 15010-1001.11	DL__Ba	<i>Hind</i> III	238 ^b	168	36864	26
<i>D. repleta</i>	REP	DSSC# 15084-1611.10	DR__Ba	<i>Hind</i> III	167 ^b	143	36864	31.6
<i>D. mercatorum</i>	REP	DSSC #15082-1521.36	DM__Ba	<i>Hind</i> III	128 ^a	125	18432	18
<i>D. mojavensis</i>	REP	DSSC # 15081-1352.22	DM_CBa	<i>Bam</i> HI	152 ^a	143	30720	28.9
<i>D. arizonae</i>	REP	DSSC# 15081-1271.27	DA_CBa	<i>Hind</i> III	152 ^c	133	18432	16.1
<i>D. hydei</i>	REP	DSSC# 15085-1641.58	DH__Ba	<i>Hind</i> III	164 ^a	146	36864	32.8
<i>D. grimshawi</i>	HAW	DSSC# 15287-2541.00	DG__Ba	<i>Hind</i> III	231 ^a	127	18432	10.1
<i>D. albomicans</i>	IMM	DSSC# 15112-1751.08	DA_BBa	<i>Hind</i> III	299 ^c	130	18432	8
<i>D. busckii</i>	DOR	DSSC# 13000-0081.31	DB__Ba	<i>Hind</i> III	194 ^b	166	18432	15.8

^aGenome size measured by PI method (Bosco *et al.* 2007)

^bGenome size measured by DAPI method a (Bosco *et al.* 2007)

^cGenome sizes of *D. arizonae* and *D. albomicans* were adopted from the genome size of a close relatives, *D. mojavensis* and *D. immigrans*, respectively.

^dCalculated genome coverage: by insert size, genome size and no of clones in the library

^eMEL : melanogaster; OBS: obscura; WIL: willistoni; VIR: virilis; REP: repleta; HAW: Hawaiian; IMM: immigrans; DOR: subgenus *Dorsilopha*

^f DSSC: *Drosophila* Species Stock Center

TABLE 2 A comparison of genomic redundancies of each *Drosophila* BAC library as estimated by empirical, hybridization and by FPC approaches.

Species	Calculated Genome Coverage ^a	Average Hyb Coverage ^b	FPC-General ^c	Ratio of a:b:c
<i>D. simulans</i>	18.2	25.0	17	1 : 1.4 : 0.94
<i>D. sechellia</i>	15.4	20.2	14	1 : 1.3 : 0.88
<i>D. yakuba</i>	9.1	11.0	9	1 : 1.2 : 1.01
<i>D. erecta</i>	18.9	19.7	14	1 : 1.0 : 0.75
<i>D. ananassae</i>	25.4	25.3	22	1 : 1.0 : 0.87
<i>D. persimilis</i>	15.2	18.3	13	1 : 1.2 : 0.86
<i>D. willistoni</i>	13.4	9.6	7	1 : 0.7 : 0.52
<i>D. americana</i>	5.7	9.1	8	1 : 1.6 : 1.36
<i>D. novamexicana</i>	8.5	14.8	13	1 : 1.7 : 1.48
<i>D. virilis</i>	17.4	32.7	19	1 : 1.9 : 1.11
<i>D. littoralis</i>	26	25.1	18	1 : 1.0 : 0.71
<i>D. repleta</i>	31.6	35.7	14	1 : 1.1 : 0.44
<i>D. mercatorum</i>	18	11.7	10	1 : 0.6 : 0.54
<i>D. mojavensis</i>	28.9	31.1	17	1 : 1.1 : 0.59
<i>D. arizonae</i>	16.1	20.2	10	1 : 1.3 : 0.63
<i>D. hydei</i>	32.8	42.9	37	1 : 1.3 : 1.12
<i>D. grimshawi</i>	10.1	14.2	9	1 : 1.4 : 0.87
<i>D. albomicans</i>	8	28.4	10	1 : 3.6 : 1.22
<i>D. busckii</i>	15.8	28.2	9	1 : 1.8 : 0.58

^aTheoretical coverage of each *Drosophila* library from the Table1

^bAverage hybridization coverage; total number of clones detected by two rounds of hybridization divided by the total number of loci; from Table S3

^cFPC based estimate of genomic redundancy of each *Drosophila* library. Total number clones in each FPC assembly divided by the total number of contigs; from Tables S4 and S5.