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March, 2007

# The Carnegie Protein Trap Library: A Versatile Tool for Drosophila Developmental Studies

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### The Carnegie Protein Trap Library: A Versatile Tool for Drosophila Developmental Studies

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> Manuscript received September 18, 2006 Accepted for publication December 18, 2006

#### ABSTRACT

Metazoan physiology depends on intricate patterns of gene expression that remain poorly known. Using transposon mutagenesis in Drosophila, we constructed a library of 7404 protein trap and enhancer trap lines, the Carnegie collection, to facilitate gene expression mapping at single-cell resolution. By sequencing the genomic insertion sites, determining splicing patterns downstream of the enhanced green fluorescent protein (EGFP) exon, and analyzing expression patterns in the ovary and salivary gland, we found that 600–900 different genes are trapped in our collection. A core set of 244 lines trapped different identifiable protein isoforms, while insertions likely to act as GFP-enhancer traps were found in 256 additional genes. At least 8 novel genes were also identified. Our results demonstrate that the Carnegie collection will be useful as a discovery tool in diverse areas of cell and developmental biology and suggest new strategies for greatly increasing the coverage of the Drosophila proteome with protein trap insertions.

THE central challenge of postsequence genomics is L to learn how an enhanced knowledge of genes, transcripts, and proteins can be applied to better understand the biology of multicellular organisms. Gaining an accurate picture of where and when metazoan genes are expressed remains a prerequisite for many such advances (STATHOPOULOS and LEVINE 2005). The discovery of distinctive, regulated programs of gene expression at a fine scale has the potential to reveal new cell types and substructures that make up tissues and the biological processes that govern their function. However, sensitive and widely applicable methods will be required to detect and distinguish developmentally programmed gene expression changes from those caused simply by cell cycling or environmental perturbation.

Several methods for analyzing gene expression within tissues are currently available. Particular cell types can sometimes be cultured *in vitro* into populations of useful size. However, isolated cells in artificial media frequently behave differently from cells *in vivo* interacting with precisely positioned neighbors in three-dimensional microenvironments. Another approach is to isolate tissue cells by flow sorting, microdissection, or laser capture and then determine their expression profiles in depth (reviewed in ESPINA *et al.* 2006). Visualizing patterns of gene expression within the intact tissues of transgenic organisms containing gene expression reporters may be the most general method (TOMANCAK *et al.* 2002). Epitope tagging, enhancer trapping, and gene trapping all have the added advantage that gene expression can subsequently be observed in living tissues, revealing dynamic processes that are largely beyond the reach of methods based on fixed material (reviewed in HERSCHMAN 2003; DIRKS and TANKE 2006).

Protein trapping is a variation of gene trapping in which endogenous genes are engineered to produce under normal controls protein segments fused to a reporter such as GFP. The great potential of this technology has been extensively documented in yeast, where large collections of strains that each trap a different gene have been generated using transposable elements (Ross-MACDONALD et al. 1999) or by homologous recombination (HuH et al. 2003). Extensive gene and protein trapping has also been carried out in cultured embryonic stem (ES) cells (Gossler et al. 1989; FRIEDRICH and SORIANO 1991), where fusions with more than half of annotated mouse genes have been recovered (see Skarnes et al. 2004). However, relatively few of these ES cell lines have so far been used to generate corresponding mouse strains where the versatility and sensitivity of the method for analyzing tissue structure can be tested.

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Large-scale protein trap screens may also reveal new information about genome structure and function. Identifying in an unbiased manner locations throughout a genome where a coding exon can be expressed tests the accuracy and completeness of its annotation. Characterizing the splicing patterns that lead to normal or aberrant GFP expression tests the current catalog of transcript isoforms generated by alternative splicing. Moreover, by recovering insertions in the same gene that splice differently and produce GFP with varying efficiency, such a project might generate a data set useful for studying the determinants of splice site selection and transcript stability.

Drosophila provides a favorable system for applying gene traps to diverse developmental and genomic studies. The genome sequence has been extensively annotated on the basis of experimental data (MISRA et al. 2002). Thousands of enhancer trap lines have been generated in large-scale transposon screens and culled of redundant strains by the gene disruption project (see BELLEN et al. 2004). In contrast, producing Drosophila protein trap lines has remained difficult. Several hundred such lines were generated using a mobile GFPcontaining exon flanked by both splice acceptor and donor sites (MORIN et al. 2001; CLYNE et al. 2003). However, the process was highly inefficient, with as few as 1 in 1500 progeny flies expressing GFP. Positive lines often contained more than one insertion, preferentially tagged a small number of hotspot loci, and tagged many sites not predicated to fuse the GFP exon in frame to any known coding region (MORIN et al. 2001). KELSO et al. (2004) found that the recovery of lines could be increased by using an automated embryo sorter to select GFP-positive embryos and established a website, Fly-Trap, to gather information on Drosophila protein trap lines. Consequently, we initiated a large-scale protein trap screen to increase gene coverage, test the genome annotation, and address some of the remaining technical difficulties in efficient line production.

Here we report the production of lines that trap 600– 900 Drosophila genes, including 244 where one or more trapped proteins can currently be identified. Using the Drosophila ovary as a test system we confirm that protein trap lines reveal fine-scale details of developmentally regulated protein expression, making them exceptionally valuable discovery tools for a wide range of studies. Finally, mapping RNA splicing patterns downstream from >1200 insertions provides insight into how an added exon affects splicing and suggests how the production of protein trap lines can be expanded to cover a larger fraction of the Drosophila proteome.

#### MATERIALS AND METHODS

**Generation of P-element lines for protein trap screening:** The P-element-based protein trap screens presented here utilized the pPG<u>A</u>, pPG<u>B</u>, and pPG<u>C</u> vectors described in MORIN *et al.* (2001). These elements carry a mini-white transgene in the opposite orientation to an enhanced green gluorescent protein (EGFP) exon, which is composed of EGFP sequence, without start or stop codons, flanked by splice acceptor and donor sites from the Drosophila MHC locus. <u>A</u>, <u>B</u>, and <u>C</u> refer to the position of the splice sites within the first and last codons of the EGFP exon sequence. Previously used pPG<u>A</u>, pPG<u>B</u>, and pPG<u>C</u> third chromosome insertions (MORIN *et al.* 2001) were remobilized in the presence of balancer chromosomes. New insertions that mapped to the *CyO* balancer chromosome, did not express EGFP, and exhibited remobilization rates off of the *CyO* balancer of at least 60% in single-pair mating assays were recovered and used as starting stocks in the screen (see below).

*piggyBac* protein trap vectors: To make a shuttle vector for subcloning the EGFP exon into different transposable elements, the entire EGFP exons from the pPG<u>A</u>, pPG<u>B</u>, or pPG<u>C</u> plasmids were excised from the original *P*-element plasmids (kind gift of W. Chia) using *Eco*RI and *Pst*I and subcloned into pBluescript (Stratagene, La Jolla, CA). These plasmids were then cut with *Eco*RV and *Kpn*I, end filled using Klenow, and religated to themselves to create pBS-GFP<u>A</u>, pBS-GFP<u>B</u>, and pBS-GFP<u>C</u>. The resulting plasmids carry the EGFP exon sequence between a unique *Eco*RI site at the 5' end and unique *Pst*I, *Sma*I, *Bam*HI, and *Xba*I sites at the 3' end. New tagging sequences can be inserted between the splice acceptor and donor sites of the exon using unique *Nco*I and *Xho*I sites.

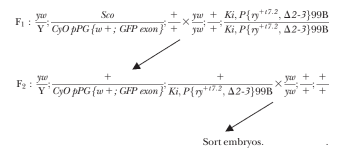
Two different *piggyBac* protein trap vectors (Figure 1) were constructed using pBac{D. m.  $w^+$ } (HANDLER and HARRELL 1999) (kind gift of A. Handler). The pBac{D. m.  $w^+$ } plasmid was digested with ClaI to remove most of the mini-white sequence and a linker containing HpaI, XhoI, and SpeI sites was inserted in its place to form pBAC{ $\Delta ClaI$ }. To create pBAC{Bg/II-GFP}, the EGFP exons from pBS-GFPA, pBS-GFPB, and pBS-GFPC were subcloned into the Bg/II and MfeI sites of pBAC{ $\Delta ClaI$ }. To create pBAC{*Hpa*I-GFP}, EGFP exon sequences were inserted between the MfeI and HpaI sites of pBAC{ $\Delta ClaI$ }. An intronless yellow transgene from the yellow-BSX plasmid (BELLEN et al. 2004) was then subcloned into the unique Spel site of both pBAC{BglII-GFP} and pBAC{HpaI-GFP} to form either pBAC{BglII-GFP;  $y^+$ }, which has the EGFP exon and yellow transgene oriented away from each other, or pBAC{*Hpa*I-GFP; y<sup>+</sup>}, which has the EGFP exon and yellow transgene pointing toward each other (Figure 1A). Both pBAC{*Bgl*II-GFP;  $y^+$ } and pBAC{*Hpa*I-GFP;  $y^+$ } vectors carrying the EGFP exon in the A frame were transformed into y w flies using the phspBac helper plasmid (HANDLER and HARRELL 1999) (kind gift from A. Handler).

We created stable genomic sources of the *piggyBac* tranposase using *P*-element transformation vectors. To place the *piggyBac* transposase under control of the ubiquitin promoter, the *piggyBac* transposase ORF was excised from phspBac using *Bam*HI and *Dra*I and ligated into the *Bam*HI and *Sma*I sites of the pCasper3-Up2-RX poly(A) *P*-element vector (WARD *et al.* 1998) (kind gift of R. Fehon), which carried a modified multiple cloning site (kind gift of A. Hudson), to form pP{UbpBACtrans}. To make an inducible *piggyBac* transposase source, phspBac was digested with *Eco*RI and *Dra*I and the fragment containing both the Drosophila *hsp70* promoter and *piggyBac* transposase ORF was ligated into the *Eco*RI and *Stu*I sites of pCasper4 to form pP{*hsp70*-pBACtrans}. These vectors were used to transform *y w* flies, using standard *P*-element transformation techniques.

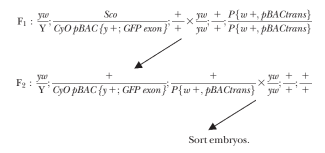
To test the activity of the *piggyBac* transposase transgenes, single-pair matings were set up using the pBAC{*HpaI*;  $y^+$ }24.3 insertion, which mapped to the X chromosome, and pP{Ub-pBACtrans} or pP{*hsp70*-pBACtrans} stocks. The pBAC{*HpaI*;  $y^+$ }24.3 insertion was mobilized in males that were then outcrossed to y w females. Phenotypically *yellow*+ males in

the next generation were scored as new insertions. The pP{*hsp70*-pBACtrans} was able to remobilize the pBAC{*Hpa*I;  $y^+$ }24.3 insert in 43% (n = 30) of single-pair matings tested whereas the pP{Ub-pBACtrans} was able to remobilize the pBAC{*Hpa*I;  $y^+$ }24.3 insert in 48% (n = 21) of single-pair matings tested. The pBAC{*Hpa*I;  $y^+$ }24.3 insertion was remobilized in the presence of a *CyO* balancer chromosome. New insertions that did not express EGFP and mapped to the *CyO* chromosome were used in the pBAC-based protein trap screen.

**Generation of embryos with novel transpositions:** We isolated new EGFP-expressing *P*-element insertions using the following genetic scheme:



New pBAC insertions were generated through a similar genetic scheme:



Hereafter, *P*-element and *piggyBac* element-based protein trap lines were treated the same. For the  $F_1$  cross several hundred males and females of the appropriate genotypes were mated in bottles to produce several thousand males in which the elements were mobilized for the  $F_2$  cross. These males were crossed to 8000–10,000 virgin *y* wfemales in a population cage. These virgin females were obtained using a virgining stock that carried a heat-shock-inducible *hid* transgene on the *Y* chromosome (kind gift of R. Lehmann). Two separate overnight embryo collections from each population cage were screened for EGFP expression. We limited the number of times we screened embryos from a particular cage to try to minimize the number of identical insertions recovered due to premeoitic insertion events.

**Embryo sorting and line establishment:** We screened for EGFP expression in embryos using a COPAS Drosophila embryo sorter (Union Biometrica). Embryos were dechorionated in 50% bleach for 2.5 min and washed extensively with water. Dechorionated embryos were then washed into sorting solution ( $0.5 \times$  PBS, 2% Tween-20). With the exception of the sorting solution, the COPAS sorter was used according to the manufacturer's protocol, using the manufacturer's solutions. The sorter and sample pressures of the COPAS machine and embryo density were maintained so that the COPAS sorter screened 15–20 embryos/sec. The sorter used a 488, 514 nm multiline argon laser. EGFP fluorescence was detected using PMT1 set to 510 nm. Red fluorescence, used as a measure of embryo autofluorescence, was detected using a second PMT set to 580 nm. Baseline values for each fluorescent axis were set

empirically using previously isolated fly strains that express low levels of EGFP and yw non-EGFP expressing embryos (Figure 1). Approximately 250,000 embryos were sorted in five 50,000embryo batches per day. Sorted embryos were collected and washed in dH<sub>2</sub>O. All the embryos from a single batch were placed together in standard food vials. We estimate that ~80% of the sorted embryos survived to adulthood. Sorted flies that survived to adulthood and did not carry the *Ki*,  $P/ry^+t7.2$ ;  $\Delta 2-3/99B$  or  $P\{w^+; pBACtrans\}$  chromosomes were individually outcrossed to a yw stock. New lines that carried EGFPexpressing insertions that did not map to the starting *CyO* chromosome were maintained as stocks.

**DNA sequencing, RT-thermal asymmetric interlaced PCR analyses, and prediction of fusion potential:** Genomic sequences flanking either *P*-element or *piggyBac* protein trap insertions were determined by members of the Lawrence Berkeley Lab group using an established protocol for sequencing inverse PCR products from genomic DNA (BELLEN *et al.* 2004). Database software developed for the annotation of the Drosophila gene disruption project (BELLEN *et al.* 2004) was used to manage the sequence data. Once the insertion site of a given protein trap line was determined, a FileMaker Pro database that contained information [version 3.2 of the Drosophila genome annotation (MISRA *et al.* 2002)] for all Drosophila transcripts, exons and introns, and their reading frames was used to predict which gene(s) and transcript(s) were being trapped by a given protein trap insertion.

We developed a reverse transcriptase coupled thermal asymmetric interlaced PCR (RT-TAIL) protocol largely on the basis of methods used to determine T-DNA insertion sites in Arabidopsis (SINGER and BURKE 2003). This method allowed us to determine the mRNA sequence adjacent to the EGFP exon without using gene-specific primers. Total RNA was isolated from 15 adult flies using an RNAqueous-96 automated kit (catalog no. 1812; Ambion, Austin, TX). The samples were ground in 200 µl of sample buffer and spun for 5 min at 14,000 rpm. The supernatant was placed in a 96-well plate and 100 µl of 100% EtOH was mixed with each sample. The sample was transferred to the filter plate, washed, and then treated with Dnase I (Ambion) for 15 min. Rebinding buffer was added to each well of the filter plate, and the plate was washed extensively. The RNA was eluted off the filter and precipitated with 7.5 M LiCl solution (Ambion). The resulting RNA pellet was washed with 75% EtOH and then retreated with Dnase I for 30 min at 37°. Dnase inactivation reagent (Ambion) was added to the samples. The RNA samples were spun and the supernatant was transferred to a new plate. A detailed protocol is available upon request.

The following GFP-specific primers were used for RT–TAIL PCR:

GFP-For1, 5'-GGAGGACGGCAACATCCTGG-3'; GFP-For2, 5'-CAACGTCTATATCATGGCCG-3'; GFP-For3, 5'-AGACCCCAACGAGAAGCGCG-3'; GFP-Rev1, 5'-GTCGTGCTGCTTCATGTGGTCG-3'; GFP-Rev2, 5'-GACACGCTGAACTTGTGGCCG-3'; GFP-Rev3, 5'-AGCTCCTCGCCCTTGCTCACC-3'.

The arbitrary degenerate (AD) primers used in this study were originally described by SINGER and BURKE (2003) but are listed here for convenience:

AD3, 5'-AGWGNAGWANCAWAGG-3'; AD4, 5'-STTGNTASTNCTNTGC-3'; AD5, 5'-NTCGASTWTSGWGTT-3'; AD6, 5'-WGTGNAGWANCANAGA-3'.

A pool of the AD primers was mixed according to <u>SINGER</u> and BURKE (2003).

The first round of RT-TAIL PCR was set up in 96-well format using a one-step RT-PCR kit (QIAGEN, Valencia, CA). For every reaction 5  $\mu$ l of total RNA was mixed with 10  $\mu$ l 5 $\times$  buffer, 2 µl 10 mм dNTP solution, 1 µl GFP-For1 or -Rev1 primer, 12.5  $\mu$ l AD primer mix, 2  $\mu$ l enzyme mix, and 17.5  $\mu$ l of dH<sub>2</sub>O. The reverse transcription reaction was carried out at 50° for 30 min. The sample was then heated to 95° for 15 min and then cycled for primary TAIL-PCR according to SINGER and BURKE (2003), using a MJ Research (Watertown, MA) thermal cycler. The secondary and tertiary TAIL-PCR reactions were carried out according to SINGER and BURKE (2003), using GFP-For2 or -Rev2 primers and GFP-For3 or -Rev3 primers, respectively, and regular TAQ DNA polyermase (Roche, Indianapolis). The PCR products of the tertiary reaction were treated with exoSAP (United States Biochemical, Cleveland) and sequenced using GFP-For3 or -Rev3 primers.

The RT–TAIL PCR protocol using the three GFP-For primers, which amplified off of the 3' end of EGFP, consistently yielded better results than the same reaction using the Rev primers. Therefore most of the RT–TAIL PCR data define splicing products at the 3' end of the EGFP sequence. To identify sequence fusing to the 5' end of EGFP, we employed a 5' RACE kit according to the manufacturer's protocol (Ambion), using the EGFP reverse primers listed above.

Analysis of protein expression in tissues: Samples were dissected in Grace's medium, placed in 48-well plates outfitted with a nylon mesh bottom, and fixed in 4% paraformaldehyde buffered in 1× PBS for 10 min at room temperature. The plate was washed extensively with PBT (1× PBS, 0.5% Triton X-100, 0.3% BSA) and incubated overnight at 4° with rabbit anti-GFP antibody (Torrey Pines) (1:2000) in PBT. The samples were then washed extensively with PBT and incubated with goat anti-rabbit Alexa488 (Molecular Probes, Eugene, OR) (1:400) for 4 hr at room temperature. The samples were then washed with 2  $\mu$ g/ml of DAPI, and mounted in Vectashield (Vector Laboratories, Burlingame, CA). Images were collected using a Leica SP2 confocal microscope.

#### RESULTS

Generating a large initial collection of tagged strains expressing EGFP: Our initial strategy was to generate a much larger number of lines containing new protein trap vector insertions than in previous screens and to institute additional technical improvements. Because of their proven utility, we used the same P-element-based protein trap vectors employed by MORIN et al. (2001), but we also constructed a similar set of vectors with piggyBac (Figure 1A). As described in MATERIALS AND METHODS, we set up crosses in small population cages to limit the recovery of clusters, utilized dominant markers to remove the transposase source from all new lines, and identified rare GFP-expressing embryos rapidly and sensitively using an automated embryo sorter (Figure 1B). This protocol allowed us to screen >60 million embryos over a period of 2.5 years, to identify >7500"green" embryos, and to use each one to start an individual culture (see Table 1). Ultimately, 7404 strains were successfully established, maintained by selection for white+ eye color, and analyzed further as diagrammed in Figure 1C.

The same scheme was used with both transposons; however, in practice the *piggyBac* vectors were not nearly

as efficient at generating EGFP-positive candidate lines as the *P*-element vectors (Table 1). *P*-element vectors typically exhibited 70% mobilization and generated ~1 EGFP-expressing embryo per 1000 sorted. In comparison, the *piggyBac* vectors displayed nearly 50% mobilization, but they yielded only 1 EGFP-expressing embryo per 50,000 sorted. Thus, the *piggyBac* vectors were slightly less efficient at mobilization, but drastically less efficient at generating EGFP-positive lines upon insertion. Consequently, we soon abandoned attempts to generate large numbers of *piggyBac* protein trap insertions (Table 1), but continued to characterize the lines we did recover to learn if they would shed any light on the lower frequency of trapping observed.

Localizing insertions on the annotated genome: To identify candidate proteins that may have been fused within individual lines, we determined the genomic DNA sequence flanking the insertion(s) in collaboration with the Berkeley Drosophila Genome Project (BDGP) gene disruption project (MATERIALS AND METHODS). In most cases, the sequences from both the 5' and 3' vector end junctions mapped by BLAST analysis to a unique insertion site within the Drosophila genome sequence. Lines for which the sequencing reaction failed, the sequence matched repetitive DNA, or the 5' and 3' sequences differed (indicating that two or more insertions were present in the stock) were recycled back into the starting pool, and frequently a unique single insert was eventually identified. Altogether the insertions in 1375 C frame, 3172 B frame, and 1009 A frame P-element and 164 piggyBac A frame lines were localized to unique genomic sites.

Knowing the genomic location of an insertion allowed us to predict which transcripts would incorporate the EGFP exon and whether they would undergo splicing and translation into a functional fusion protein. First, we removed ~1550 duplicate lines derived from premeiotic clusters that were identified because they bore insertions identical in position and orientation to those in one or more sibling lines. Of the 4170 independent lines remaining, 2149 (52%) were associated with a gene correctly oriented for possible fusion (*i.e.*, located between -500 and the 3' end). We also classified the ways an insert can be located relative to its closest annotated transcript into general categories as diagrammed in Figure 1D and classified all the lines (Table 2).

Expression of a fusion protein is expected when the GFP exon resides between two coding exons within an intron of matching reading frame (class 1A). Such insertions made up only 23% of the total localized insertions and defined 192 different genes (Table 4). Forty percent of insertions were close to an annotated gene but were not predicted to express the EGFP exon (classes 2–4), while 37% of the lines were not located within 0.5 kb of a correctly oriented gene. EGFP production from these lines might be explained by the use of unannotated

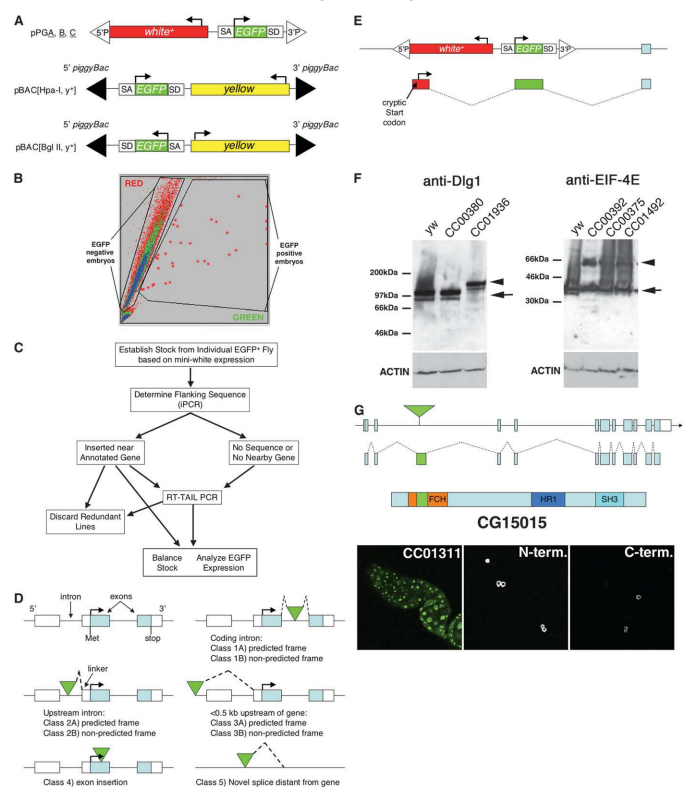


FIGURE 1.—Generation and classification of protein trap vector insertions. (A) Schematic of protein trap vectors (after MORIN *et al.* 2001). (B) Sample output from automated sorting of Drosophila embryos mobilized from site not expressing GFP. Rare GFP+ embryos (red circles) registering above a threshold value are diverted by the machine and later used to start individual cultures. (C) Scheme for characterization of putative protein trap lines (see text). (D) Classification of the general types of relationships between transposon inserts and the local genome annotation. Classes 1–4 consist of insertions in the appropriate orientation located within a codon intron (class 1), a noncoding transcribed region (class 2), an upstream genomic region (class 3), or an exon (class 4). For each class, the insert was either of the appropriate frame (subclass A) or of nonappropriate frame *(Continued)* 

1510

TABLE 1 Project summary

Туре	No.	%
Total setup	7404	100
CA	1344	18
CB	4000	54
CC	1870	25
piggyBac A	190	3
Aligned sequence	5720	77
Clusters	1550	21
Independent aligned	4170	56
Gene hits	2149	29
Spacing hits	2093	28
Different genes	1154	16
Protein traps	244	
Enhancer traps	256	
Novel gene/exon	50	
Unclassified	300	
Balanced stocks	878	12

TABLE 2

Line types

		Lines	
Туре	Code	N	%
Coding intron, in frame	1A	1337	23
Coding intron, out of frame	1B	173	3
Noncoding intron, in frame	2A	29	1
Noncoding intron, out of frame	2B	429	8
1-500 bp upstream, in frame	3A	56	1
1–500 bp upstream, out of frame	3B	756	13
Exon	4	804	14
>500 bp to next oriented gene	5	2093	37
Total		5677	100

See class definitions in Figure 1D.

spliced into the EGFP exon from the 5' P-element sequences of the vector. The P-element promoter is highly efficient at enhancer trapping, and the entire first exon of the transposase gene is present in the vector along with the start of intron 1, which is in the frame compatible with CB lines. These lines were associated with nuclear localized EGFP, possibly due to fusion of the first exon of *P* transposase with EGFP. Further evidence of enhancer trapping was observed in the analysis of line CA07138. The EGFP RNA was fused to sequences, including an in-frame ATG start codon, derived by transcription and splicing from the noncoding strand of the mini-white transgene carried in the P-element vector (Figure 1E). These observations suggest that EGFP expression in a significant number of the lines depends on transcripts initiated from within the transposon itself by enhancer trapping rather than on EGFP exon addition by splicing into exogenous transcripts.

Analysis of downstream transcript sequences: To gain additional information, we analyzed the sequences downstream from the EGFP exon from many lines in the collection by carrying out RT–TAIL–PCR in a 96-well format (MATERIALS AND METHODS). 3' sequences up to 700 bp in length and defining the location of one to six downstream exons were obtained for >1200 lines (Table 3). The pattern of downstream splicing allowed productive fusions to be identified and indicated lines that splice out-of-frame and likely become subject to nonsense-mediated decay (NMD) (VASUDEVAN and PELTZ 2003). Fusions within lines of classes 2–4 often

genic elements, by noncanonical splicing events, or by the presence of a second insertion at a canonical site.

The Drosophila genome annotation is highly supported by experimental evidence (MISRA et al. 2002); hence the frequency of these discrepancies was surprising, but a similar outcome has been reported in previous protein trap screens using both yeast (Ross-MACDONALD et al. 1999) and Drosophila (MORIN et al. 2001). Some protein-coding genes may have been missed within cDNA libraries (HILD et al. 2003), and a substantial number of genes may contain unannotated far-upstream promoters and alternative translation start sites. There might be a large class of RNA genes that have escaped detection but that can drive expression of EGFP using cryptic start sites. Alternatively, the high selective pressure used to isolate EGFP+ strains may have led to the recovery of rare events in which the normal gene or transcript structure has changed.

**Analysis of fusion transcripts:** We sequenced portions of the fusion transcripts to address how EGFP expression arises in lines of various classes. A limited number of 5' RNA sequences were obtained by 5' RACE analysis or by TAIL–PCR (see MATERIALS AND METHODS). As expected, several lines in class 1A were found to initiate in normal exons upstream from the insertion site. However, we discovered several CB lines that

<sup>(</sup>subclass B) to fuse to the protein if splicing continued to the next annotated exon splice acceptor site. Class 5 consists of transposons inserted >0.5 kb from a correctly oriented annotated gene. (E) The structure of cryptic transcripts initiated within the Drosophila mini-white marker gene that contain an ATG codon and splice in frame to EGFP, thereby allowing expression independent of an endogenous transcript in some lines. (F) Western blot analysis of Dlg1 and eIF-4E protein production in control animals (*y w*, CC00380) and insertion lines predicted to trap Dlg1 (CC01936) or eIF-4E (CC00392, CC00375, and CC01492). (G) Abnormal nuclear accumulation of CG15015-EGFP in line CC01311 whose insertion lies within the FHC domain (left). Tissue culture cells expressing N-terminal or C-terminal fusions are found in the cytoplasm (center and right).

TABLE 3 RNA analysis

Туре	Successful	Confirmed DNA	Second insert	% confirmed
CA	316	224	50	82
CB	328	166	75	69
CC	572	165	72	70
Piggy A	13			
Totals	1229	555	197	74

were predicted to encode a "linker peptide," which might or might not include a stop codon, derived from the translation of a small segment of upstream nucleotides. The RNA analysis also revealed the presence of a second insertion in 14% of type 1A lines, but between 24 and 50% of the other classes. The second insertions found within class 2–5 lines were often valid protein trap alleles (class 1A) and were frequently the true source of the lines' EGFP production. This information allowed us to identify additional candidate fusions (Table 4), to correct many initial line classifications, and to more accurately estimate the total number of trapped genes (Table 1: 600-900). By the time lines were selected and balanced, secondary insertions or damage did not contribute substantially to the phenotypes reported in Table 4. Tests estimated the frequency of background lethal mutations among balanced, saved lines at 7-21%, similar to the best transposon screens (SPRADLING et al. 1999).

Novel splicing suggests new genes and exons: We compared the splicing observed downstream from the inserted exons with that of the genome annotation (MISRA et al. 2002) to identify new Drosophila gene and transcript isoform candidates. To identify new candidate genes, we focused on lines inserted >0.5 kb from an appropriately oriented known gene and for which RNA sequence data were also available. In 114 of these 205 lines, the RNA sequence coincided with the position and orientation of the insertion and therefore indicated the splicing pattern downstream of the single EGFP exon. Most of the lines spliced to one or more novel exons. At least 8 probably correspond to unannotated genes because they match previous gene predictions (HILD et al. 2003) or are supported by EST data (see Table 4). Most of the remaining exons do not predict proteins with homologs in other species and represent either aberrant splicing events or novel or untranslated exons.

Similar analysis of 297 lines with intron insertions allowed us to test for novel exons and transcript isoforms. We examined 443 splicing events and identified a total of 35 (7.9%) that did not correspond to current gene models (MISRA *et al.* 2002). Since at least some of these differences probably resulted from aberrant splicing induced by the insertions, this represents a maximum estimate of the fraction of unannotated genomic exons and emphasizes the high accuracy and completeness of current Drosophila gene models, at least for abundant transcript forms. Often, the RNA data indicated which isoform among several predicted to fuse in frame is likely to predominate in ovarian tissue. For example, we could determine that line CA06613 in ovarian tissue predominantly fuses the Su(Tpl) gene rather than *Mi-2*, in whose transcription unit it also lies in frame.

The nature of the noncanonical splices observed was interesting. The most common events (21/35) were for insertions in large introns to splice to a novel exon(s) prior to joining the predicted downstream exon. Some simply appear to define alternative isoforms that skip exons or utilize different exon combinations not previously documented. Some of these events may have been induced by the abnormal position of the EGFP exon within the primary transcript. However, several lines appear to define alternative isoforms because they utilize different combinations of known exons in no previously documented transcript isoforms. Three lines utilized 5' start sites for exons that differed by 6, 21, or 27 bp from the annotated exon. The CC01473 transcript reads through an annotated exon into the adjacent intron and probably defines a novel alternate transcript 3' end. Although we consider it likely that many of these differences reflect endogenous Drosophila gene expression, all of the candidate novel genes and transcript isoforms require independent confirmation in strains lacking protein trap insertions. Such tests were beyond the scope of our project.

Protein trap insertions likely vary in the fraction of the endogenous protein that is tagged with EGFP for a variety of reasons. First, in many lines only some of the multiple-transcript isoforms contributing to protein production are tagged by the insertion and fused in frame. Second, the splicing efficiency of the EGFP exon might vary due to its surrounding genomic context. To investigate this issue, we analyzed the protein products of tagged genes by Western blotting. The tagged proteins were easily distinguished from their wild-type counterparts on the basis of size and by probing with protein-specific and anti-EGFP antibodies (Figure 1F). In line CC01936 all three isoforms are predicted to incorporate the EGFP exon in frame, and nearly all of the ovarian Dlg1 protein incorporated EGFP as indicated by its mobility. A similar result was reported previously in the case of line CB02119 (BUSZCZAK and SPRADLING 2006), where the precursors of five of six annotated transcripts are predicted to contain the insertion, although only two fuse in frame. In contrast, only  $\sim 50\%$  of the ovarian wild-type eIF-4E protein is tagged with EGFP (Figure 1F) despite the fact that six of seven annotated eIF-4E transcripts initiate upstream

	proteins
<b>TABLE 4</b>	d trapped
	Identified

		2002		Strand	rnenotype	11	Insert	Met	dore	12	IIISCII	Met	otop	T3	Insert	Met	Stop	$\mathbf{T4}$	Insert	Met	stop	ıype
•	CA06506	14068962	3R	+	Lethal	RA	1.5	1	4	RB	1.5	1	4	RC	1.5	1	4	RD	1.5	1	4	1A
26-29-p C	CA06735	13987313	3L	+	hv	RA	0.5	1	3													3A
-	CC00758	21169155	2L	+	Lethal	RB	1.5	1	4													1A
Actn C	CC01961	1927332	X	Ι	hv	RB	2.5	0	10	RA	2.5	5	10	RC	2.5	5	10					1A
AGO1 C	CA06914	9841633	2R	Ι	hv	RC	3.5	00	x	RA	3.5	5	1	RB	0.5	5	1					1A
Alh C	CC01367	2935653	3R	Ι	hv	RA	5.5	3	10	RD	5.5	ы	10									1A
apt C	CC01392	19468319	2R	+	hv	RB	1.5	1	Ŋ	RD	1.5	ы	9	RE	1.5	3	9	RC	1.5	1	ъ	1A
-	CC01186	19473808	2R	+	hv	RB	1.5	1	Ŋ	RD	2.5	0	9	RE	2.5	3	9	RC	1.5	1	ъ	1A
	CB03579	417662	X	+	hv	RA	3.5	1	Ŋ													1A
k	CB05492	9051633	3L	Ι	Lethal	RA	3.5	0	4	RB	2.5	0	60									1A
	CB03789	9056078	3L	I	Lethal	RA	2.5	n cv	4	RB	0.5	n cv	60									1A
-	CC00319	16783418	3R	+	Lethal	RC	1.5	\$	10	RA	1.5	1	10	RB	0.5	64	6	RD	0.5	\$	9	1A
-	CC01941	17072582	X	+	hv	RA	1.5	1	1													1A
-	CC00869	4485350	3R	Ι	hv	RA	1.5	1	4													1A
t1	CB02354	5996196	3R	Ι	hv	RA	1	3	1													4
56D	CC02069	15338563	2R	Ι	Lethal	RB	1.5	1	0	RC	1.5	2	0	RD	1.5	0	0					1A
	CB02667	16418866	3R	+	Semilethal	RA	0.5	1	10													3B
Bsg C	CA06978	8104393	2L	+	hv	RB	2.5	0	1	RA	2.5	0	1	RD	2.5	5	1	RC	2.5	0	4	1A
-	CB03431	12482943	2L	I	hv	RA	3.5	-	ъ	RB	1	-	30	RD	1.5	1	3	RE	2.5	5	4	1A
Cam	CC00814	8149270	2R	+	Lethal	RA	2.5	0	ы													1A
CAP C	CA06924	6190378	2R	+	hv	RI	9.5	1	14	RH	7.5	ŝ	12	RJ	8.5	3	13	RG	7.5	3	12	1A
CAP C	CA07185	Transposon	2R	+	hv	RI/RF	3.5	1	14													1A
Cat C	CC00907	18815951	3L	+	Lethal	RA	1.5	1	3													1A
-	CC01469	10063184	2R	+	hv	RD	1.5	1	10	RB	1.5	Г	11	RC	2.5	3	11	RA	2.5	5	11	1A
10724	CA07499	13406231	3L	+	hv	RA	1.5	1	1	RB	1.5	1	1									1A
-	CA06844	12472570	×	I	hv	RC	1.5	1	4													1A
-	CB04917	13015302	3L	I	hv	RA	1.5	1	3													1A
-	CC01391	7033157	2L	+	hv	RA	2.5	2	9	RD	2.5	×	6	RG	2.5	7	œ	RF	1.5	9	1	1A
-	CC06238	4764704	3R	+	Semilethal	RA	2.5	2	6													1A
Ŭ	CC00625	1076935	3R	I	hv	RA	1.5	-	9	RB	1.5	Γ	5 CI									1A
-	CC06135	12098103	3R	I	Lethal	RA	1	-	9													4
-	CC01646	1650510	3L	+	Lethal	RA	1.5	1	3													1A
-	CB02069	19502492	X	+	Lethal	RA	1.5	1	4	RB	1.5	1	5 C									$^{1A}$
CG1440 C	CA07287	8331699	X	+	hv	RA	1.5	1	Ŋ													1A
CG14648 C	CA06610	229231	3R	+	hv	RA	1.5	1	9	RB	1.5	3	9									1A
CG14656 C	CA06996	624015	3R	Ι	Lethal	RA	2.5	1	3													1A
CG15926 C	CB04063	12365107	X	+	hv	RA	0.5	0	ы													3A
-	CB03410	3416244	2R	Ι	hv	RA	1.5	0	3	RC	1.5	1	3	RB	1.5	0	3					1A
-	CC01294	16654986	3R	I	Lethal	RA	1.5	1	Ŋ													1A
CG17646 C	CB02833	1737431	2L	+	hv	RB	1.5	61	12	RA	0.5	5	12									2B
CG1888 C	CB02075	5434043	2R	Ι	hv	RA	1.5	1	5													1B

p Type	1A	IA	1A	1A	1A	1A	1A	1A	1A	1A	1A	2B	2B	1A	1A	1A	2A	1A	1A	1A	1A	1A	4	1A	1A	1A	3A	1A	3A	1A	1A	1A	$1\mathrm{A}$	3A	$1\mathrm{A}$	1A	1A	1A	1A	1A	$1\mathrm{A}$
Stop							x								Ŋ			5 D				5															5				
Met							3								3			1				0															Γ				
Insert							1								1.5			1.5				2.5															1.5				
T4							RC								RD			RH				RG															RE				
Stop	4						10						3		5 2			9		5		x		4	5 2	5 2										4	9	5			4
Met	6						%						1		1			1		1		0		5	2	1										5	5	1			Γ
Insert	0.5						3.5						0.5		1.5			1.5		2.5		2.5		1.5	1.5	1.5										1.5	1.5	0.5			2.5
T3	RD						RB						RA		RB			RA		RB		RA		RA	RB	RC										RA	RC	RC			RC
Stop	4			60		14	6				4	5 2	4		Ŋ		4	9		5	2	6	3	3	4	9										8	ъ	2		30	4
Met	1			1		0	0				1	2	2		2		6	1		1	1	1	0	1	2	2										61	0	1		1	2
Insert	1.5			1.5		2.5	2.5				1.5	1.5	1		1.5		1	1.5		2.5	0.5	3.5	0.5	1.5	1.5	2.5										2.5	1.5	0.5		1.5	2.5
T2	RA			RD		RD	RD				RC	RB	RB		RC		RB	RI		RC	RA	RF	RB	RC	RA	RB										RB	RD	RA		RA	RA
Stop	4	4	9	9	ы	13	10	10	8	x	ы	9	4	3	ы	1	4	9	8	Ŋ	9	6	4	0	4	ы	0	4	3	3	×	ы	1	1	4	×	Ŋ	5	4	4	4
Met	6	6	-	1	1	0	%	0	1	1	5	3	3	1	1	1	6	1	5	1	5	1	3	1	1	1	3	1	1	1	1	1	1	П	1	61	5	1	1	0	6
Insert	1.5	2.5	1.5	4.5	1.5	2.5	3.5	4.5	1.5	3.5	2.5	2.5	1.5	1.5	1.5	1.5	1.5	1.5	2.5	2.5	2.5	3.5	6	1.5	1.5	0.5	0.5	1.5	0.5	1.5	1.5	4.5	1.5	0.5	1.5	2.5	1.5	1.5	1.5	2.5	95
$\mathbf{T1}$	RB	RA	RC	RC	RA	RB	RA	RA	RD	RA	RB	RA	RC	RA	RA	RA	RA	RI	RA	RA	RB	RE	RA	RB	RC	RA	RA	RA	RB	RA	RA	RA	RA	RA	RA	RC	RB	RB	RA	RB	RB
Phenotype	v	Λ	Λ	Λ	Λ	Λ	hv	Lethal	hv	Λ	hv	Lethal	Λ	Λ	hv	Lethal	hv	hv		Lethal	Lethal	Lethal	hv	hv	Lethal	hv	Lethal	hv	Λ	Λ		Lethal	v	Λ	v	Λ	Λ	Λ	Λ	Λ	Λ
	hv			hv	hv .	hv	h	Γ	h	hv		Γ	hv	hv	h		Ч			Γ	L	Γ	Ч	h	Γ	h	Γ	h	hv	hv			hv	· hv	hv	hv	hv	hv .	. hv	hv	hv
Strand	+	+	+	+	I	+	I	+		+	+	+	I	+	I	+		+	I	Ι	Ι	+	I	I	+	I	I		+	+	I	+	+	I	+	+	+	I	I	+	+
Chr	3R			3R	2L	3L	3L	3L	3L	X	04	3R	X	X	3L	3R	3L	2R	3L	3L		3L	2R	3R	3L	3R	3L	3L		3L	2R	3R	2R	2L	3L	2R	2L	3R	3R	2L	2R
Site <sup>a</sup>	27573862	4894157	26640622	26646037	2873276	10515608	5177892	882041	3060297	17632469	2699967	5176796	1802294	3069871	20509742	16079041	20486552	14309292	15001227	15808775	22779020	8627040	13604099	7392313	15099355	7597410	15088263	8308089	17647286	7969693	4863309	4631299	12075198	8159580	7354388	16208199	20824559	163496	9226998	2755189	9852057
Line	CC01491	CA06801	CA06686	CA06810	CA07748	CC00511	CC00236	CA06614	CC00904	CA06772	CB04483	CC01586	CA07694	CA07562	CA06926	CC00526	CB03619	CA07176	BA00207	CA07529	CB03223	CC00858	CC00677	CA06960	CA07332	CA06556	CC00864	CC00645	CB04101	CB02086	BA00169	CA07228	CA06604	CA07352	CA06603	CC00825	CB04962	CB02188	CC00817	CC00719	CC01377
Gene	CG1910	CG3036	CG31012	CG31012	CG31694	CG32062	CG32423	CG32479	CG32486	CG32560	CG3287	CG33936	CG3810	CG3939	CG5059	CG5060	CG5130	CG5174	CG5392	CG6151	CG6330	CG6416	CG6424	CG6783	CG6854	CG6930	CG6945	CG7185	CG7484	CG8209	CG8213	CG8351	CG8443	CG8552	CG8583	CG8920	CG9331	CG9772	CG9796	CG9894	Cp1

# TABLE 4 (Continued)

Gene																						
	Line	Site <sup>a</sup>	Chr	Strand	Phenotype	IT	Insert	Met	Stop	T2	Insert	Met	Stop	T3	Insert	Met	Stop	T4	Insert	Met	Stop	Type
J	CA06507	5456158	3R	I	hv	RA	2.5	1	4													1A
J	CB03073	16280297	2L	Ι	hv	RA	1.5	1	60													1A
CycB C	CC01846	18694008	2R	Ι	hv	RA	1.5	Г	5	RD	1.5	1	ы	RB	1	1	5 C	RC	0.5	1	4	1A
	CC00921	12743940	2R	+	hv	RA	1.5	1	11	RC	1.5	Г	11	RB	0.5	Г	10	RD	1.5	1	6	1A
J	CA06616	12744777	2R	+	hv	RA	2.5	1	11	RC	2.5	1	11	RB	1.5	Г	10	RD	2.5	1	6	1A
desat1 C	CC01694	8269738	3R	+	hv	RA	1.5	2	5	RC	1.5	0	Ŋ	RE	1.5	2	Ŋ	RB	1.5	0	ы	2B
C	CB02104	21629393	2L	Ι	hv	RB	2.5	ы	4	RA	1.5	1	0	RF	1.5	1	0					1A
Ō	CC01936	11286274	X	+	hv	RF	5.5	1	6	RB	6.5	3	17	RH	5.5	2	16	RE	2.5	3	13	1A
J	CA06573	6480860	2L	+	hv	RA	0.5	1	4													3A
C	CB03889	24717483	3R	+	hv	RA	3.5	1	13	RB	3.5	5	13	RE	0.5	0	12	RF	3.5	1	13	1A
B	BA00164	17211471	2R	+		RD	1.5	5	15	RA	1.5	0	14	RE	1.5	5	11					2A
J	CA06594	9111298	2R	+	hv	RA	1.5	-	6													1A
Ū	CC00251	19190343	2L	+	hv	RA	0.5	Г	4													3B
Ũ	CC01924	21681694	2L	Ι	Lethal	RA	1.5	1	5 C	RC	1.5	0	ы	RB	1	5	Ŋ					1A
Ũ	CC01915	10565092	3R	Ι	Lethal	RA	2.5	5	9													1A
eIF-2β C	CC06208	12519527	3L	Ι	Lethal	RA	1.5	1	3													1A
eIF3-S9 C	CB04769	13423919	2R	+	hv	RB	1.5	2	5	RA	1	1	4									2A
eIF-4a C	CB03721	5982474	2L	+	Lethal	RA	1.5	1	5	RC	1.5	1	Ŋ	RB	0.5	1	Ŋ	RD	0.5	1	Ŋ	1A
eIF-4E C	CC00392	9395078	3L	Ι	hv	RD	1.5	2	9	RB	1.5	0	9	RC	1.5	1	5 C	RA	1.5	0	9	1A
eIF-5A B <sub>2</sub>	BA00155	19945688	2R	+		RB	1.5	2	4	RA	1.5	0	4									2A
eIF-5C B <sub>2</sub>	BA00280	1425192	3R	Ι		RA	1.5	2	8	RC	2.5	3	6	RF	1.5	5	8	RD	1.5	0	8	2A
Eip63E C.	CA06742	3569399	3L	+	hv	RD	2.5	Ч	11	RE	3.5	0	12	RA	4.5	3	12	RB	3.5	0	11	1A
J	CA06515	12435789	2L	+	hv	RA	1.5	1	1													1A
Ŭ	CB03931	20342600	3L	+	Lethal	RC	2.5	2	x	RB	2.5	0	x	RE	2.5	2	x	RG	2.5	5	x	1A
Q	CB03613	4029454	X	Ι	hv	RA	6	2	10													4A
-	CC01359	16403817	3L	I	Lethal	RA	1.5	-	Ŋ	RC	1.5	-	ъ С									lA
_	CA06503	26212791	3R	I	Lethal	RA	1.5	Γ	0	RB	2.5	21	4	RC	2.5	2	4	RD	2.5	21	4	$\mathbf{IA}$
Fer2LCH C	CA07607	26215006	3R	+	Lethal	RA	с, ,	<i></i> , 0	4	RB	с, ,	<i></i> 60	4	RC	1	- ,	21					4,
	CC01493	17185787	X	I	hv	KA	C.1	-	C I	KC	2.5	N	9	KU	0.5	Τ	D D					IA
Fkbp13 C	CA07340	17385064	2R	Ι	hv	RA	0.5	2	5	RB	1.5	Г	2 2									IA
-	CB04937	7194698	2 <b>R</b>	Ι	hv	KA	1.5	-	9													IA
Fs(2)Ket C	CA07301	20735659	2L	+	hv	RA	2.5	21	9													lA
J	CA07108	9493967	2L	I	Lethal	RA	1.5	-	0													1A
-	CB02804	12106314	3R	+	hv	RD	2.5	2	12	RB	2.5	0	12	RE	2.5	0	12	RA	0.5	3	12	$^{1A}$
GlcAT-S C	CA07168	9616795	2L	+	hv	RA	1.5	1	5 C	RB	0.5	1	IJ									1A
-	CB02989	15762784	2L	Ι	hv	RA	0.5	5	1	RB	0.5	2	Ŋ	RC	0.5	3	x	RD	0.5	2	1	3A
A	CA06658	6331783	2R	+	Semilethal	RB	2.5	5	x	RC	2.5	60	6	RD	2.5	2	x	RE	2.5	5	x	1A
-	CC00195	1647884	2R	+	hv	RA	0.5	1	19													Ŋ
HDAC4 C	CA07134	13172850	X	Ι	hv	RA	2.5	5	14	RC	1.5	1	12									1A
-	CC00664	27763272	3R	Ι	Lethal	RB	4.5	4	14	RA	4.5	4	14	RK	3.5	\$	13	RH	5.5	5 C	16	1A
His2Av C	0000320	99603903	9 D	+	I athal	Δ	с 7	-	~													14

(Continued)

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Type	1A	1A	1A	1A	1A	1A	1A	1A	1B	1A	1A	1A	3A	1A	1A	1A	1A	1A	1A	1A	1A	2A	1A	2A	1A	2B	$1\mathrm{A}$	1A	IA	IA ;	IA	IA	$\mathbf{IA}$	$1\mathrm{A}$	1A	1A	1A	1A	1A	1A	1A
Stop				Ŋ	5		9				9										13			9											Ŋ						
Met				1	Г		3				0										1			1											51						
Insert				0.5	2.5		2.5				0.5										1			1.5											4.5						
T4				RC	RC		RD				RE										$\mathbb{RF}$			RD											RD						
Stop	7	8		Ŋ	5 C	Ŋ	9	6			50										19			9									ŋ		Ŋ					3	
Met	1	0		1	1	1	0	3			Γ										3			4									ß		3					1	
Insert	1.5	2.5		0.5	2.5	1.5	2.5	0.5			1.5										7.5			1.5									1.5		4.5					1	
T3	RC	RC		RB	RB	RC	RG	RA			RA										RJ			RA									RB		RF					RA	
Stop	9	1	3	Ŋ	5	9	4	6			ъ	3		1	1		9		4		19			9				9	16	ſ	Ū,	4	ß	4	Ŋ	6				60	14
Met	1	0	1	1	1	2	0	3			1	1		1	1		5		1		0			3				2	-	c	N ·		ß	1	2	5				1	1
Insert	0.5	2.5	1.5	1.5	2.5	2.5	3.5	1.5			1.5	0.5		1.5	1		0.5		1.5		7.5			1.5				2.5	1.5	1 ,	C.1	2.5	1.5	1.5	4.5	7.5				1.5	1.5
12	RB	RB	RB	RE	RE	RB	RH	RC			RH	RB		RB	RB		RB		RA		RB			RC				RA	RB	6	KB	RA	RA	RD	RA	RB				RB	RB
stop	9	x	3	Ŋ	5	9	9	6	ы	0	4	0	6	18	18	12	1	2	5 C	4	19	4	4	9	4	Ŋ	2	1-	16	<b>~</b> `	Ū,	4	ß	4	Ŋ	6	6	10	60	3	12
Met	1	0	1	1	П	2	2	1	0	1	1	1	1	3	3	5	1	1	2	1	0	5	1	4	1	5	П	<i></i>	_		_		-	-	5	2	3	Ι	1	3	1
Insert	1.5	2.5	1.5	1.5	2.5	2.5	2.5	1.5	4.5	1.5	1.5	1.5	0.5	12.5	12.5	4.5	2.5	1.5	2.5	2.5	7.5	1.5	1.5	1.5	1.5	1.5	1.5	3.5	1.5	1.5	c.1	2.5	1.5	1.5	4.5	7.5	3.5	1.5	2.5	1.5	1.5
11	RA	RA	RA	RA	RA	RA	RF	RB	RB	RA	RD	RA	RA	RA	RA	$\mathbf{RA}$	RA	RA	RB	RA	RL	RA	RA	RB	RA	RA	RA	RB	RA	RA	KA	RB	RA	RC	RR	RA	RA	RA	RA	RA	RA
Phenotype	hv	Lethal	hv	hv	Lethal	Lethal	hv	hv	hv	hv	hv	hv	hv	hv	hv	hv	hv	hv	hv	Lethal	Lethal	Lethal	Lethal	hv	hv	hv	hv	hv	hv	hv	hv	hv	Lethal	hv	hv	hv	hv	hv	hv	hv	hv
Strand	I	+	I	+	+	+	Ι	I	I	+	Ι	+	I	I	I	Ι	+	Ι	Ι	+	I	+	I	I	I	+	I	I	+	+ -	+	Ι	I	Ι	I	+	I	+	+	I	+
Chr	2L	3R	3R	3R	3R	3L	X	3L	X	3R	3R	3R	2L	2L	2L	X	X	X	2R	3L	3R	2L	2R	3R	X	3L	3R	2R	3R	3R	ZL	3R	3L	3L	3R	3L	X	X	3R	3R	3L
Site	6722933	17881934	9485980	24425969	24427038	14031299	10702676	18833638	6892590	6182245	7430651	25608214	12822834	220632	221924	19525602	15393000	9447368	14688632	1336285	1123169	5543070	10462132	24152038	14969607	21502447	27891553	13756894	20060119	22978192	10240237	8545125	19901556	11546502	17200382	21909824	14596460	13534781	25831370	4873684	12141797
Line	CB02121	CA07414	CC00189	CC01563	CA06921	CB02656	CB04573	CC00377	CB04539	CC00294	CB05190	CC01156	CB02190	CC01466	CC00801	CC01368	CC00492	CA06684	CA06962	CA07460	CA07520	CB03749	CB04957	CC06230	CA07051	CA06602	CC00109	CC01398	CC00924	CB04968	CB05282	CC06325	CA06598	CB04396	CA07012	CC01995	BA00253	CA07788	CC01224	CB02647	CA06597
Gene	homer	how	Hrb87F	Hrb98DE	Hrb98DE	Hsc70Cb	Imp	Indy	inx7	jumu	Jupiter	kay	kekl	kis	kis	1(1)G0084	1(1)G0168	1(1)G0320	1(2)08717	1(3)02640	1(3)82Fd	Lam	LamC	larp	Lsd-2	M6	Map205	Mapmodulin	mask	Mdh	me31B	Men	Mi-2	Mob1	mod(mdg4)	mub	NetB	NFAT	NIp	Nmdmc	Nrx-IV

TABLE 4 (Continued)

<b>TABLE 4</b>	(Continued)

Type	1A	1A	3A	1A	1A	1A	1A	1A	1B	1A	1A	1A	4A	1A	1A	1A	1A	1A	1A	1A	$1\mathrm{A}$	3A	3A	3A	1A	1A	1A	1A	1A	4	$^{1A}$	$^{1A}$	lA	1A	1A	1A	1A	4	1A	1A	$1\mathrm{A}$
Stop T								5			10						9	11			9				5 C											10					
Met S								1			5						4	1			5				1	1										5					
Insert								0.5			2.5						2.5	6.5			1.5				1.5	0.5										2.5					
T4 I								RC			RD						RB	RB			RB				RA	RA										$\mathbf{RF}$					
Stop							18	ы	4		10						5 D	13			9	6			9	9						9			15	10					
Met							6	3	5		3						1	0			0	1			5	0						0			0	5					
IIISCII							0.5	1.5	1.5		2.5						2.5	8.5			2.5	0.5			0.5	2.5						2.5			2.5	2.5					
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done			4	16	ъ	3	20	61	9		11		00			10	8	13	ы		7	3			9	9		9			10	1			15	10	0		14	6	4
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1125111			0.5	4.5	2.5	1	4.5	1.5	1.5		2.5		1			1.5	3.5	8.5	2.5		2.5	0.5			2.5	2.5		1			1.5	2.5			2.5	2.5	2.5		2.5	2.5	1
11			RA	RA	RB	RB	RE	RB	RB		RC		RA			RC	RD	RD	RB		RC	RC			RE	RE		RB			RA	RA			RB	RA	RA		RA	RC	RA
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INICI	1	1	1	0	0	1	0	1	1	0	3	2	1	1	0	0	4	0	1	1	5	1	Г	3	0	0	1	1	5	2	0	2	Г	1	0	61	1	1	1	3	2
TIDSTIT	2.5	1.5	0.5	4.5	2.5	1.5	3.5	1.5	1.5	2.5	2.5	2.5	1	1.5	3.5?	2.5	3.5	8.5	1.5	1.5	1.5	0.5	0.5	0.5	3.5	3.5	1.5	1.5	4.5	10	3.5	2.5	2.5	1.5	2.5	2.5	1.5	1	0.5	2.5	2.5
11	RA	RA	RA	RB	RA	RA	RA	RA	RF	RA	RB	RA	RB	RA	RB	RB	RC	RA	RA	RA	RE	RA	RB	RB	RB	RB	RA	RA	RA	RC	RB	RC	RA	RA	RC	RB	RB	RA	RC	RA	RB
rnenotype	Lethal	hv	hv	Lethal	hv	hv	hv	Lethal	hv	Lethal	hv	hv	hv	Lethal	Lethal	hv	hv	Lethal	Lethal	Lethal	Mfsterile	Lethal	hv	Lethal	hv	hv	hv	hv	Lethal	hv	hv	hv	Lethal	hv	hv	hv	Lethal	Lethal	hv	hv	hv
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	3L	2R	3R	3R	2R	3R	2R	3L	3L	2R	2R	3R	2R	3L	2R	X	3L	3R	3R	2R	3R	3L	3R	2L	2L	2L	3L	3R	3R	3R	X	2R	2L	2L	2R	X	3L	3L	3L	2R	2L
OILC	Transposon	8060372	9789604	13529472	4019736	8523601	19554469	15134669	7847944	12548787	5157995	19712573	4996761	20303643	Transposon	11538719	1501291	4983771	16937950	2584592	1833559	11815918	1449810	19009261	5000772	4997815	5802962	26714539	18184952	22393784	15712098	17362946	6479544	8592539	12411885	2536739	7361764	21024944	2107593	15504045	1119094
TILLC	CC01311	CB03751	CB04400	CC00445	CC00380	CB02132	CA07101	CA06526	CB02246	CA07474	CC01654	CB03040	CB02099	CC01326	CC01645	CC06344	CA06961	CC00479	CA07717	CA07465	CB02119	CB02653	CC01920	CB03373	CA06523	CA06547	CC01583	CA07241	CA07674	CA07683	CA07575	CC00871	CC00735	CA07125	CA06989	CA06683	CC01823	CC01921	CA06744	CC00233	CA07211
Gene	Oda	Oda	Orc2	osa	Pabp2	Past1	Pde8	Pdi	Pdp1	Picot	Pkn	Pli	Pmm45A	polo	bsd	Ptp10D	pÙf68	und	Rab11	Rab2	Rm62	RpL10Ab	RpL13A	RpL30	Rtnl1	Rtnl1	S6k	Sap-r	sarl	scrib	sd	Sdc	$Sec61\alpha$	Sema-la	Sema-2a	Sgg	$Sh3\beta$	Sin	sls	sm	smi21F

Type	1A	3A	4A	3A	1A	1A	1A	1A	1A	1A	1A	1A	1A	1A	1A	1A	1A	4	1A	1A	1A	1B	1A	1A	1A	1A	1A	1A	1A	1A	1A	1A	$1\mathrm{A}$	1A	1A	1A	1A	1A	1A	5B	5B
Stop								8			x		10	10	1	1		4								4				4					0						
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Stop						Ŋ		6			9		10	10	1	1		9							16	×				60					3						
Met						1		1			1		0	0	3	3		3							3	1				1					3						
Insert						1.5		1.5			1.5		3.5	3.5	2.5	6.5		2.5							2.5	5.5				1.5					2.5						
T3						RA		RD			RN		RD	RD	RF	RF		RA							RC	RC				RC					RB						
Stop	3			5 C		9		1	5		9	4	10	10	9	9		50		4		5			16	×		×		4	4	60			ы	0		14	9		
Met	1			2		1		1	0		1	5	6	6	1	1		3		1		1			3	5		1		5	1	1			1	1		1	3		
Insert	1.5			0.5		1.5		1.5	2.5		1.5	1.5	3.5	3.5	1.5	5.5		2.5		1		1			2.5	5.5		1.5		2.5	1.5	1.5			1.5	0.5		2.5	4.5		
T2	RA			RC		RC		RH	RB		RC	RB	R	R.	ß	RD		RB		RB		RA			RA	RE		RA		RB	RA	RA			RD	RA		RA	RB		
Stop	12	61	3	4	9	1	3	8	4	11	\$	5	10	10	1	1	x	4	4	4	ы	3	4	4	16	1	4	ы	00	4	Ŋ	4	6	6	3	3	4	14	2		
Met	1	1	5	2	1	1	1	1	1	1	0	1	0	0	2	2	1	3	1	1	0	1	1	1	2	0	1	0	1	2	2	5	ы	ы	1	1	1	5	3		
Insert	1.5	0.5	1	0.5	2.5	1.5	1.5	1.5	1.5	1.5	0	1.5	3.5	3.5	2.5	6.5	1.5	3	1.5	1.5	2.5		1.5	1.5	2.5	4.5	1.5	5	1.5	2.5	2.5	2.5	2.5	2.5	1.5	1.5	1.5	2.5	4.5		
T1	RB	RA	RA	RA	RA	RB	RA	RI	RA	RA	RB	RA	RB	RB	RE	RE	RA	RC	RA	RA	RA	RA	RA	RA	RG	RB	RA	RA	RA	RA	RB	RB	RA	RA	RA	RA	RA	RB	RA		
Phenotype	hv	Lethal	Lethal	hv	hv	hv	Lethal	hv	Lethal	hv	Semilethal	hv	Lethal	Lethal	hv	Lethal	hv	hv	Lethal	hv	hv	Lethal	Lethal	hv	hv	hv	Lethal	Lethal	Lethal	Lethal	Lethal	Lethal	Lethal	hv	hv	hv	Lethal	Lethal	hv	hv	hv
Strand	I	I	Ι	+	+	I	I	+	Ι	Ι	Ι	+	+	+	+	+	+	I	+	+	+	I	I	I	Ι	Ι	Ι	+	Ι	Ι	+	Ι	Ι	Ι	+	I	I	I	I	+	+
Chr	X	3L	2L	3L	×	3R	3L	2R	3R	X	X	2R	3R	3R	3R	3R	X	2R	3L	X	2R	3R	2R	3L	2R	3R	3L	3R	3R	2R	3R	3R	3R	2L	3L	2L	3L	2R	3L	3R	3L
$\operatorname{Site}^a$	13104608	12727708	9897503	22766451	6162464	9470746	14402679	5004591	11171291	7861288	6986034	5877016	11116123	11117364	26389239	26400579	15214506	10491357	12508905	8137659	2903327	21707093	19932991	262842	11681495	20044629	5358821	14538229	15469816	2518996	1417892	8452738	15045962	5019005	970771	6918786	12117267	20896845	16102655	4638102	8022556
Line	CC01032	CB02932	CB02294	CA06644	CA07692	CB02655	CA07249	CC02013	CC01684	CC01711	CB05562	CB04973	CC01710	CC00578	CC00416	CA07346	CC01414	CC01925	CA06517	CA06750	CC01420	CC01830	CC01393	CC00737	CC00482	CA06641	CA07496	CA06666	CA07644	CA06708	CC01380	CA07634	CB05330	CC00791	CA07004	CB03248	CA06791	CC01626	CA07703	CB02318	CC01309
Gene	sno	snRNP69D	sop	SPoCk	Spt6	sqd	stwl	Su(var)2-10	Surf4	SWS	Sxl	TER94	Tm1	Tm1	tmod	tmod	Top1	tra2	tral	Trxr-1	Tsp42Ee	Tsp96F	tsr	Tudor-SN	tun	twin	UevlA	VAChT	Vha13	Vha16	Vha26	Vha55	vib	vkg	NSG	xl6	yps	diz	Zn72D		

TABLE 4 (Continued) Protein Trap Lines in Drosophila

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Jene L	ine	Site <sup>a</sup>	Chr	Strand	Gene Line Site <sup>a</sup> Chr Strand Phenotype	T1	Insert	Met	Stop	T2	T1 Insert Met Stop T2 Insert Met Stop T3 Insert Met Stop T4 Insert Met Stop	Met	Stop	T3 j	nsert	Met	Stop	T4 I	nsert	Met	Stop	Type
CB(	02658	CB02658 12522067	2R	Ι	hv																	5B
CC	CC01670	12944837	3R	+	hv																	5B
CC	CC01932	21856862	3R	Ι	hv																	5B
CAI	CA07436	5644358	2R	Ι	sl																	5B
CB(	CB03064	8536402	2L	+	hv																	5B
CC	00523	CC00523 22806145	3R	Ι	hv																	5B

from the insertion site. These examples suggest that the EGFP incorporation level varies between genes in large part due to the tagging of a subset of gene isoforms that themselves display differing expression levels.

In contrast, we found little evidence of short-range context effects. Less than twofold variation in protein expression as measured by Western blotting with anti-EGFP antibodies was observed between lines with insertions at sites within the same intron (N. SRIVALI and A. SPRADLING, unpublished data). However, these experiments did reveal that insertions of the *piggyBac*based vector consistently produced less EGFP protein than lines with the corresponding *P*-element-based vector that were inserted in the same intron (N. SRIVALI and A. SPRADLING, unpublished data). This suggests that some aspect of the structure of the *piggyBac* vector used compromised splicing efficiency.

Identification of protein trap and enhancer trap core collections: To help identify a core set of valid gene trap lines we examined the EGFP expression patterns of many nonredundant lines in both the adult ovary and larval salivary gland. There was a strong correlation between insert location and the nature of the staining patterns observed. More than 95% of lines in class 1A, the in-frame fusions, produced patterned EGFP expression above background in at least some ovarian cells or in the salivary gland. In contrast, a much smaller, but still significant, fraction of lines in classes 2-5 also expressed EGFP in a regulated manner. Combining information on insert location, genome annotation, RNA transcript sequence, and EGFP pattern, we identified a set of 244 lines predicted to produce fusions between EGFP and 431 protein isoforms of 233 distinct genes (Table 4). These new protein trap lines express EGFP in a wide variety of cellular compartments under diverse developmental controls (see Figures 2 and 3).

A second major class of lines in the collection showed the properties expected of EGFP enhancer traps (Table 5). These CB lines were susceptible to enhancer trapping from the *P*-element promoter, were located mostly upstream of the annotated start site, expressed EGFP in nuclei, and the RNA analysis, if available, did not indicate fusion in frame downstream. The expression patterns of such insertions in well-characterized genes supported this interpretation. For example, line CB02030 in *ptc* showed strong expression in the inner sheath cells of the germarium (FORBES *et al.* 1996), while line CB04353 in *Dad* strongly expressed in the germline stem cells and immediately downstream germ cells (KAI and SPRADLING 2003; CASANUEVA and FERGUSON 2004).

The characterization of a significant number of lines in the collection remains incomplete (Table 1). Many of these contain insertions located >0.5 kb from an appropriately oriented annotated gene but where RNA sequence data were not obtained. Others are inserted within genes at locations not predicted to generate protein fusions or gene traps. Some of these lines

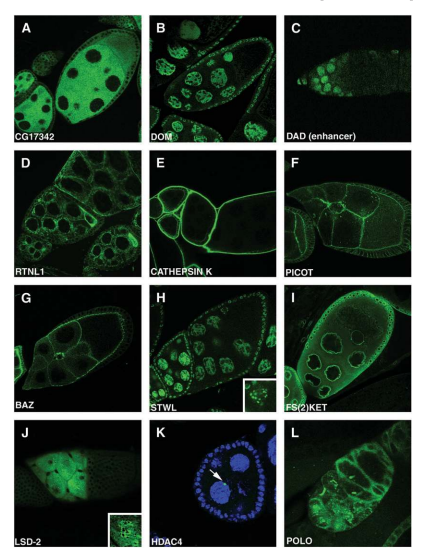


FIGURE 2.—Protein traps for the study of protein subcellular localization. Patterns of subcellular localization of EGFP expressed from the following lines that trap the indicated genes were observed: (A) cytoplasmic, CA06607 (CG17342); (B) nuclear, BA00164 (dom); (C) enhancer trap nuclear, CB04353 (Dad) in stem cells and early cystocytes; (D) endoplasmic reticulum, CA06523 (Rtnl1); (E) extracellular, CA06735 (cathepsin K); (F) membrane, CA07474 (Picot); (G) apical, CC01941 (Baz); (H) chromatin, CA07249 (stwl); (I) nuclear membrane, CA07301 (Fs(2)Ket); (J) lipid droplets, CA07051 (Lsd-2); (K) novel structure, CA07332 (CG6854); (L) novel structure, CC01326 (polo).

express EGFP in the ovary, and we cannot rule out that others express transcripts in other tissues. On the basis of the processing of previous lines in these same classes this suggests that a significant number of new enhancer traps and a handful of new protein traps could be sorted out from a larger number of lines with secondary insertions in already trapped genes. Consequently, the number of different genes trapped in the collection is likely to increase beyond the 600 or so currently characterized.

Even when a protein is tagged in frame, the insertion of the EGFP sequence is expected to disrupt its normal structure and localization some fraction of the time. For example, line CC01311 traps CG15015, the Drosophila homolog of mammalian Cip4, a modular protein that interacts with Cdc42 and helps to regulate the actin cytoskeleton (ASPENSTROM 1997). The CC01311 *P* element is inserted between the first two coding exons and thus disrupts the FES/Cip4 domain of CG15015 (Figure 1G). The protein trap fusion product localizes to the nucleolus while transgenes of CG15015 tagged at either the very N or C termini localize to the cytoplasm when expressed in S2 cells. We observed that 3 other protein trap fusion products of 107 analyzed accumulated in the nucleus when they were expected to reside in the cytoplasm.

Diverse behavior of tagged proteins: The 244 identified protein trap lines of the core collection exhibit extremely diverse patterns of EGFP expression, suggesting that proteins occupying a wide range of cellular compartments can be tagged in vivo. We observed many lines with EGFP fluorescence in the cytoplasm (Figure 2A) or nucleus (Figure 2B) as expected. Localization to intracellular membranous structures was also commonly seen, as illustrated by a trap in the ER structural component Reticulon-1 (Figure 2D). Gene trapping of secretory proteins is thought to be inefficient due to retention of the fusion proteins in the ER where the activity of the fusion gene may be affected (SKARNES et al. 1995). The full-length protein traps we constructed could label secreted proteins, as indicated by the extracellular localization of EGFP in CA06735, a fusion

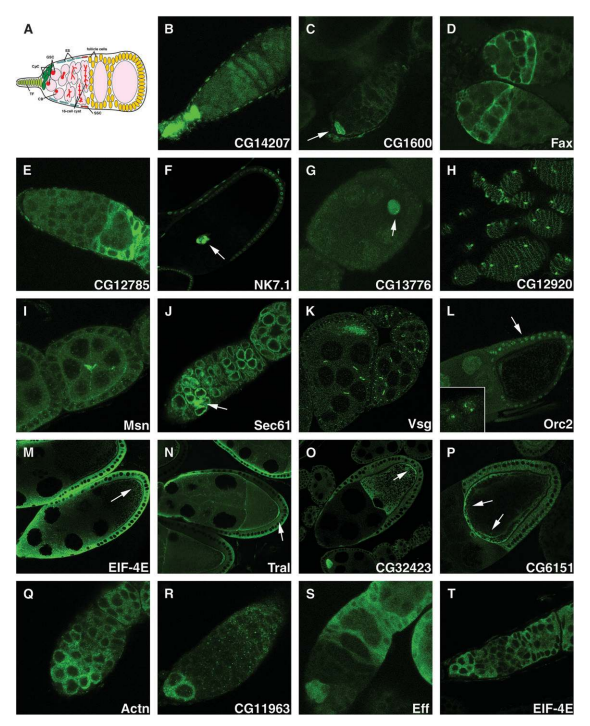


FIGURE 3.—Protein traps for the study of developmental regulation during oogenesis. The expression in the ovary of various protein trap lines is shown to illustrate how they can be used to associate genes with developmental processes. (A) Schematic of an ovariole tip. The terminal filament (TF), cap cells (CpC), germline stem cells (GSC), cystoblast (CB), and escort cells (ES) are illustrated. (B–H) Cell type identification. (B) Terminal filament, CB02069 (CG14207); (C) cap cells, CB03410 (CG1600); (D) escort cells, CC01359 (fax); (E) follicle cells, CC06135 (CG12785); (F) outer border cells and posterior follicle cells, CB02349 (NK7.1); (G) oocyte nucleus equals the germinal vesicle (arrow), CB04219 (CG13776); (H) novel sheath cell type, CC01646 (CG12920). (I–L) Analyzing developmental processes. (I) Novel structure in center of midstage follicle, CC00523 (Msn); (J) fusome, CC01436 (Sec61); (K) germline and somatic ring canals, CA07004 (Vsg); (L) chorion gene amplification, CB04400 (Orc2). (M–P) Localization of proteins in the oocyte. (M) Posterior pole, CC01442 (EIF-4E); (N) posterior pole, CA06517 (Tral); (O) posterior pole, CC00236 (CG32423); (P) anterior pole, CA07529 (CG6151). (Q–T) Developmental regulation of gene expression in early germ cells. (Q) Control with little change, CC01961 (Actn); (R) GSC/CB enriched, CC06238 (CG11963); (S) GSC and early cyst enriched, CC01915 (Eff); (T) GSC and forming cyst enriched, CC01442 (EIF-4E).

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$ \begin{array}{rrrrr} CB04422 & 6068307 & 3L & + & hv & RA & 1 \\ CB04175 & 708235 & 3L & - & hv & RA & 0.5 \\ CB02255 & 19546304 & X & + & hv & RA & 0.5 \\ CB02236 & 6879851 & X & + & hv & RA & 0.5 \\ \end{array} $	$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$		1	1												3
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CB02236 6879851 X + hv	CB02236 6879851 X + hv CB02133 13345190 2R + hv		1	7											.,	3
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TABLE 5

Enhancer trap alleles

					17				ł				T		1100111		and			- I.	and the
CG14709 C CG1621 C CG1667 C	CB02373	21878549	3R	Ι	Lethal	RA	0.5	1	9												60
	CB04397	7394886	3R	+	hv	RA	0.5	1	8												3
-	CB02042	3380602	2R	Ι	hv	RA	0.5	1	5												3
	CB05477	5731280	2R	+	Semilethal	RA	2	1	Ŋ												4
CG16708 C	CB04388	1193533	3R	Ι	hv	RA	1.5	ы	4												0
CG16817 C	CB02934	5503678	3R	+	hv	RA	1	1	4												4
CG16971 C	CB03898	224106	3L	+	hv	RB	1	ы	00	RC	1	3	3	RD	1	3	00				4
CG17002/vimar C	CB05094	2873307	2R	+	hv	RB	1	1	л С												4
CG17090 C	CB03116	544248	3L	+	hv	RB	1.5	0	10												5
CG17090 C	CB02270	544285	3L	Ι	hv	RA	1.5	1	01												5 C
-	CB02962	18823565	2L	+	hv	RA	0.5	6	5												3
CG17836 C	CB05263	14749549	3R	+	hv	RB	3.5	3	5	RA	3.5	0	5 C	RC	2.5	3	4	RD (	0.5	2	1
$1(3) \pm 1921$	CB03702	27573214	3R	Ι	Lethal	RB	1.5	0	4	RA	0.5	1	4								0
CG2051 C	CB03625	1614858	3R	+	hv	RB	1	1	3	RA	0.5	1	3	RC	0.5	1	\$				4
CG2186 C	CB05020	10751743	X	Ι	hv	RA	1	1	1												4
CG2446 C	CB03703	11608741	X	Ι	hv	RA	0.5	8	4	RE	1	3	4	RB	1	8	4	RD	_	2 3	3
CG2698 C	CB03026	3827319	3R	+	hv	RA	1	0	x												4
CG2865 C	CB03023	2187549	X	Ι	hv	RA	0.5	1	0												3
CG2926 C	CB02232	1414090	3R	+	hv	RA	0.5	1	1												3
CG2974 C	CB04047	9980614	X	Ι	hv	RA	0.5	1	0												60
CG30055 C	CB02739	8477121	2R	+	hv	RA	1	1	1												4
CG30497 C	CB02106	3667177	$2\mathbf{R}$	Ι	hv	RA	0	0	3												4
CG31241 C	CB02140	14081632	3R	+	Lethal	RA	0	0	5												4
CG31475 C	CB04600	15006356	3R	+	hv	RA	0.5	0	4												3
CG31522 C	CB02693	279018	3R	Ι	hv	RB	1	ы	10	RC	1	ы	5								4
CG3164 C	CB02987	129446	2L	+	hv	RA	1.5	1	8	RC	1.5	1	x	RB	0.5	1	1	RD (	0.5	1 7	1
CG31650 C	CB05467	5043131	2L	Ι	hv	RA	1.5	3	4	RB	1	3	4	RC	1	3	4				3
	CB03570	20429045	2L	+	hv	$\mathbb{R}A$	1	0	9												4
-	CB03239	2735300	2L	+	hv	RC	1.5	0	11	RD	1.5	0	11	RA	1.5	0	11	RB	5.	2 11	61
-	CB03345	16716141	2L	Ι	hv	RA	2.5	1	9	RB	2.5	1	7								1
3	CB03247	9455668	3L	+	hv	RB	5	5	ũ	RA	5	0	ũ								4
CG3209 C	CB05445	19956511	2R	+	hv	RA	1	1	1	RB	1	1	9								4
CG32345 C	CB03988	628140	3L	Ι	hv	RA	0.5	1	1												5 C
CG32436 C	CB02614	21340125	3L	Ι	hv	RA	1.5	1	5 2												1
CG32436 C	CB04148	21340268	3L	+	hv	RA	1.5	1	л С												1
CG32486 C	CB03414	3070840	3L	+	Lethal	RD	0.5	1	8												3
CG3321 C	CB05150	10153408	3R	+	hv	RA	61	0	01	RB	61	6	0								3
CG33214 C	CB04173	21500637	3L	Ι	hv	RA	1	1	9												4
CG33232 C	CB03740	2466875	3L	+	hv	RA	0.5	3	6												3
CG33558 C	CB05689	2859128	2R	Ι	hv	RA	0.5	6	16												3
CG33967 C	CB04401	10549498	3R	I	Lethal	RA	1	1	6												4

		2002	IIIO	סוומווט	rnenotype	11	112011	INTAL	and	-	1176111	MICI	dore	C1	THSELL	IMEL	dore	14	TIDSTIL	INTEL	done
CG33982	CB04965	15527849	3L	+	hv	RA	1	1	1												
CG3409	CB04412	2602327	2R	I	hv	RA	1.5	3	7												
CG34110	CB04263	21010670	3R	+	Lethal	RC	2.5	Г	6												
CG3428	CB02131	9480857	3L	Ι	hv	RA	1	1	3												
CG3654/Uch-L3	CB04163	9470143	3L	Ι	hv	RA	0.5	Г	4												
CG4091	CB02284	19589843	2R	Ι	Lethal	RA	1	3	3	RC	1	3	4								
CG4300	CB04249	12270328	3L	Ι	hv	RA	1	1	4	RB	1	1	4								
CG4570	CB02124	6682617	3R	+	hv	RA	1	-	1												
CG4612	CB05331	20413641	2R	Ι	hv	RA	0.5	3	3												
CG5381	CB04616	10321949	2L	+	hv	RA	0.5	3	7												
CG5543	CB04854	19696764	2R	+	hv	RA	0.5	Г	1												
CG5548	CB05667	14957872	X	Ι	hv	RA	0.5	0	0												
CG5677	CB02054	20055723	3R	Ι	Lethal	RA	1	-	1												
CG6014	CB04785	21390388	3L	+	hv	RA	1.5	3	7												
CG6218	CB03213	11168745	3R	+	hv	RA	1	3	5 C												
CG6311/Nedd4	CB04145	17522938	3L	+	hv	RC	1	-	ъ												
CG6439	CB03836	17844824	3R	Ι	Lethal	RA	1	1	9												
CG6499	CB05192	11075733	3R	I	Lethal	RA	5	Ч	ъ												
CG6540	CB03922	1854228	X	+	hv	RA	1	Г	6												
CG6770	CB02632	12046132	2L	Ι	hv	RA	0.5	1	1												
CG7110	CB04925	13399559	2L	+	hv	RB	5	3	7												
CG7228	CB03115	7994181	2L	I	hv	RA	1	3	IJ	RB	1	3	5 L								
CG7331	CB05154	4175563	3R	+	hv	RA	1	Г	1												
CG7637	CB03644	6698443	2R	I	hv	RA	0.5	Ч	61												
CG8036	CB04958	4495764	3R	+	Lethal	RB	2.5	3	4	RC	1.5	3	\$								
CG8092	CB05336	11101113	2R	Ι	Lethal	RA	1	1	9	RB	1	-	3								
CG8128	CB02087	15591543	X	+	hv	RA	1	3	4												
CG8206	CB04527	15634596	X	+	hv	RA	0.5	-	1												
CG8444	CB03837	5086366	3R	+	Lethal	RA	1	Ч	3												
CG8583	CB04752	7353717	3L	I	hv	RA	1	-	4												
CG9062	CB02056	7171458	2R	I	hv	RB	1	Γ	x												
CG9171	CB02706	5800187	2L	I	hv	RA	1	4	x	RB	1	0	1								
CG9328	CB03031	20797707	2L	Ι	hv	RB	0.5	51	0												
CG9591	CB05694	9508902	3R	Ι	Lethal	RA	2	51	6												
CG9666	CB04503	19257579	3L	I	Lethal	RA	1	3	4												
CG9699	CB02196	16581750	Χ	Ι	hv	RA	2.5	3	ы	RF	2.5	0	Ŋ	RB	2.5	2	ъ	RD	2.5	0	5 C
CG9821	CB03335	4646100	3R	Ι	Lethal	RB	1	5	5	RA	1	0	0								
CG9921	CB02890	16226793	X	Ι	hv	RA	0.5	3	3												
CG9924	CB03517	9852398	3R	Ι	hv	RB	2.5	3	6												
Chd64	CB03690	4122396	3L	Ι	hv	RB	1	Γ	3												
chrb	CB05429	11480949	3L	+	hv	RC	1	Г	4												

TABLE 5 (Continued)

TABLE 5	(Continued)

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7110	11100985	11809411	22260750	8837940	20696520	15803675	12882290	8820577	5599791	21142183	19439096	19748795	0040400	00/6070	06616161	14299509	9967309	9389656	11268107	16172362	14561061	1426796	959579	3460609	18007641	749995	14732350	1727775	15333865	9510510	14120268	9510094	20085050	3632190	2136127	18758843	21298814	11898010	19163698	17681968	17087979	11988609
711117	CB04933	CB03039	CB02217	CB04073	CB02981	CB02618	CB04353	CB03495	CB05233	CB04282	CB03160	CB09988		CD02103	CB02040	CB03754	CB02226	CB02974	CB04090	CB02620	CB04040	CB02125	CB05493	CB05358	CB05160	CB09035	CB05084	CB02039	CB02017	CB02992	CB04634	CB05447	CB05794	CB02956	CB02790	CB05043	CB03489	CB05224	CB02997	CB02364	CB05177	CB05139
Actic	Cp190	crol	Csp	CtBP	CycB3	CycD	Dad	dally	dap	Dap160	Dcn-1 / nita	Dely 1/ puu	Jane 1	uesau	DI .	Dp1/imd	Dref/RpL13	drk	dup	eas	edl	eIF	eIF3-S10	elF5B	Fin75R	emc	endoA	Eno	esg	fas	fj	Africal Action of the test of test	fok/neb	for	fs(1)K10/kz	ftz-fl	Furl	fw	fz2	glec	Glycogenin	Grip163

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Line	CB04894	CB04956	CB05261	CB04444	CB03117	CB05774	CB09943	CB09648	CD0202010	CB02849	CB05039	CB05603	CB03168	CB02227	CB09050	CB09000	CB09331	CB05457	CB04179	CD08178	CB031/3	CB02898	CB02120	CB02888	CB03167	CB04603	CB02150	CB02595	CB04813	CB02689	CB03632	CB04106	CB02172	CB03663	CB04377	CB04551	CB05687	CB02076	CB02349	CB02015	CB04635	CB04883	CB04897
Gene	grp	GstEl	GstS1	gukh	Gyc76C	hdc	HI.Hm7	Hman	ugiiiu	HPIb	Hr39	Hsc70-4	Hsr&ohgr	IP3K1	Irn	k117	1(9)øl	lama	L an A		LBK	lea	Lk6	lola	Map60	mbc	Mbs	MESR3	<b>MESR4</b>	mirr	Mnf	Mocs1	mod	mRpS17	Myo31DF	neb/fok	nes	Neu3	NK7.1	omu	omu	$\operatorname{Nrg}$	orb2

TABLE 5 (Continued)

Protein Trap Lines in Drosophila

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Lethal hv hv		TICHORNE TT		AT HIDCHT	MCL	dore	12	Insert	Met	stop	13	Insert	Met	Stop	T4	Insert 1	Met Stop	p Type
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		RA	A C	).5	5	12	RC	0.5	60	13	RD	0.5	0	12				3
Lethal		RA	A 1		5	x	RB	1	5	x								4
		RA	A I		1	5 C												4
		RA	A C	.5	5	4												39
Lethal		RA	A	.5	2	5 C												5
Lethal		RA	A I		1	4												4
Semilethal			В	).5	5	10												3
		RA	A		1	4												4
		$\mathbb{R}$	A 1		1	5												4
		RA	A I		1	9												4
		R	A C	.5	1	4												39
		RA	A		1	00												4
Lethal		R	A 1		-	2												4
		R	A		1	3												4
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TABLE 5 (Continued)

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Gene	Line	Site <sup>a</sup>	Chr	Strand	Site <sup>a</sup> Chr Strand Phenotype Tl Insert Met Stop T2 Insert Met Stop T3 Insert Met Stop T4 Insert Met Stop	$\mathbf{T}\mathbf{I}$	Insert	Met	Stop	T2	Insert	Met	Stop	T3	Insert	Met	Stop	T4	Insert	Met		Type
trx	CB05156	10108478	3R	I	hv	RA	1.5	3	6	RB	1.5	3	8	RC	1.5	2	7	RD	1.5	2	8	2
ttk	CB02274	27550755	3R	+	hv	RE	1.5	5	5	RF	1.5	3	5	RB	1	0	5	RC	1	5	5	5
Ugt37c1	CB02900	12733228	$2\mathbf{R}$	Ι	Lethal	RA	0.5	1	1													60
Ugt86Da	CB03314	6982675	3R	+	hv	RA	0.5	1	3													60
Vha100-2	CB03404	14224630	3R	Ι	hv	$\mathbb{RB}$	1.5	ы	9													3
VhaPPA1-1	CB02209	10729723	3R	Ι	Lethal	RA	1	3	2													4
wun2	CB02267	5301760	$2\mathbf{R}$	+	hv	RA	0.5	1	9													3
Xbp1	CB02061	17031115	$2\mathbf{R}$	+	Lethal	RB	1	-	3	$\mathbf{RA}$	1	1	3									4
XeĨ	CB05610	1495043	3R	+	Lethal	RA	1	Γ	x													4
Z4/CG12974	CB02305	21279245	3L	+	Semilethal	RA	1	1	61													4
<sup>a</sup> Annotatio	<sup><math>a</math></sup> Annotation release 5.0.	.0.																				

TABLE 5 Continued) with CG8947, a Drosophila cathepsin K homolog (Figure 2E), and the membrane location of a trap in Picot, a phosphate symporter (Figure 2F). Proteins that are apically localized in polarized epithelia such as ovarian follicle cells were easily visualized, as observed for Bazooka (Par3) (Figure 2G). Subcompartmentalized nuclear proteins were also readily apparent. For example, a trap of the Stonewall (Stwl) HMG-related protein involved in germ cell chromatin organization (CLARK and MCKEARIN 1996) labeled nurse cell nuclei and the oocyte nucleus (inset) differently (Figure 2H). Fs(2)Ket, a protein involved in nuclear import, was localized to the nuclear periphery (Figure 2I). In some cases, cell-specific cellular compartments were labeled, such as in CA07051, which traps Lsd-2 and exhibited EGFP localization to lipid droplets that arise in latestage nurse cells (Figure 2]).

These studies provide a high-resolution view of known protein locations in living cells and also identify many proteins that were not previously known to reside within these compartments. In addition, the value of protein trapping as a discovery tool was illustrated by the fact that we observed new patterns of localization as well. For example, the HDAC4 protein, fused by the CA07134 trap, labeled a small body often found in only one nurse cell within an egg chamber (Figure 2K). A spindle-like structure in young nurse cells was labeled with a protein trap in the *polo* gene encoding a mitotic kinase (Figure 2L). Antibodies specific for the trapped protein can be used to isolate and further investigate the proteins present in these novel structures.

Analysis of developmental regulation—ovarian cells: All the lines in the core collection were characterized on the basis of their patterns of expression in germ cells and follicle cells during oogenesis. These experiments identified lines expressing in the major classes of somatic cells, including terminal filament cells (Figure 3B), cap cells (Figure 3C), escort cells (Figure 3D), profollicle cells (Figure 3E), and border and posterior cells (Figure 3F). Other lines expressed in germ cells of various ages, including some that were highly enriched in the oocyte nucleus (germinal vesicle) (Figure 3G). As in the case of subcellular compartments, these studies documented patterns of developmental expression for many genes that were not previously known. These genes become attractive candidates for study of their function in the corresponding processes.

Strikingly, the collection also revealed the likely existence of new cell types and novel biological processes previously unrecognized despite many years of study of ovarian biology. Line CC01646 traps the CG12920 protein and is expressed in a small subset of ovarian sheath cells that likely represent a novel cell type (Figure 3H). In line CC00523 we observed accumulation of Msn-EGFP preferentially at the center of midstage growing follicles (Figure 3I). It was not previously known that this region was the site of unique protein accumulation. Msn encodes a protein involved in Jun kinase signaling, suggesting that a special intercellular junction may assemble in this region to structurally organize the nurse cells. We observed a similar expression program (not shown) for the line CB03040 that fuses the *Pli* gene, encoding a protein associated with the NF-κB signaling response.

Developmentally specific subcellular structures, including the fusome (Figure 3J, arrow) and both somatic and germline ring canals (Figure 3K), were also labeled by rare lines. A general and extremely useful application of the collection is to identify new proteins that are associated with such structures and analyze the effect of mutations. For example, the preferential accumulation of Sec61 in the fusome observed here has been validated in recent studies (SNAPP *et al.* 2004). Proof of principle experiments of this type that focus on the fusome will be described elsewhere.

Another valuable capacity of protein trap lines is the ability to follow important developmental processes at high resolution and in living cells. During oogenesis, at least four major clusters of chorion genes undergo specific gene amplification in stage 10B follicles, a process that can be visualized as small "amplification dots" of BrdU incorporation (CALVI et al. 1998). The amplifying genes specifically contain substantial amounts of replication initiation proteins such as Orc2 at this time, whereas normally Orc2 is found throughout the cell nuclei (ROYZMAN et al. 1999). A protein trap line in Orc2 allows the amplifying loci to be directly visualized (Figure 3L). Inspection shows that the dots are not present in preamplification stage follicle cells but strongly label amplifying gene loci at stage 10B (Figure 3L, arrow).

The Drosophila oocyte represents an important model system for studying RNA and protein localization. Several biochemical and genetic studies have identified proteins enriched at either the anterior or the posterior pole of the oocyte (LASKO 2003; WILHELM and SMIBERT 2005). Protein traps in genes identified in these studies, including EIF-4E (Figure 3M) and Tral (Figure 3N), faithfully recapitulate the localization patterns of their endogenous counterparts to the posterior pole of the oocyte (WILHELM et al. 2003, 2005). Several other proteins tagged in the collection display posterior localization patterns including a trap in CG32423 (Figure 3O), a largely uncharacterized RNA-binding protein. In addition, a trap in CG6151 appears to be enriched at the anterior of the oocyte (Figure 3P). While future work will clarify the role of these proteins in oocyte patterning, these examples show that protein trapping can complement other approaches and be used to identify new components of localized RNP complexes within the cells.

Protein traps provide unique opportunities to analyze gene regulation during development in populations of cells that are difficult to isolate and in those that are sensitive to loss of cellular context. We illustrate the potential of this approach using the regulation of germ cell development within and just downstream from the germline stem cells (GSCs). Many protein trap lines, including CC01961 in Actn, showed uniform expression in GSCs, CBs, and developing germline cysts (Figure 3Q). However, it was possible to find other examples where expression levels between GSCs and early germ cells differed from those in other germ cells within the germarium. One of the most striking examples was line CC06238 that traps the putative Drosophila succinate CoA ligase gene. Expression was stronger in stem cells (and sometimes early cystoblasts) than in later germ cells as illustrated in Figure 3R. Several other genes were downregulated shortly after GSC division, including effete (Figure 3S), encoding the UbcD1 ubiquitin-conjugating enzyme that has been shown to affect germline cyst formation (LILLY et al. 2000). Another line whose EGFP expression was downregulated slightly later, at the completion of cyst formation, trapped the Drosophila eIF-4E gene (Figure 3T). Downregulation of a related gene CG8023 was previously observed at a slightly earlier time, during cyst divisions (KAI et al. 2005).

Studies on the limitations of current protein trap methods: We also tested the sensitivity of the approach used here to detect Drosophila genes by looking at the expression of the lines in the core collection in germline stem cells. First, although lines were selected on the basis of expression in embryos, we found ovarian expression above background in >90% of lines in the core collection. However, this does not address whether many other genes exist that were fused but expressed EGFP at levels too low to detect in either tissue. Analysis of germline stem cell RNA by hybridization to Affymettrix arrays detected transcripts from ~6500 Drosophila genes over an ~1000-fold dynamic range (KAI et al. 2005). Although translational regulation and differential protein stability, not to mention differences in staining sensitivity between different preparations, would be expected to introduce potential variation, we were curious whether protein trap lines could detect stem cell gene transcripts across the full range of expression levels.

We observed a strong correspondence between these two measures of stem cell gene expression (Figure 4, A– E). Lines with very strong EGFP expression tended to have RNA levels at least 10-fold higher than lines with above background but relatively weak expression. These lines in turn had signals higher than most lines scored as below the level of detection on arrays. The correlation was not perfect; for example, some lines showed more EGFP expression in stem cells than might be expected from the microarray study (Figure 4F). The existence of such lines was not unexpected, because some lines likely still carry second insertions, and the microarray used was based on release 1 gene models. Overall, we could detect EGFP above background in GSCs from nearly all

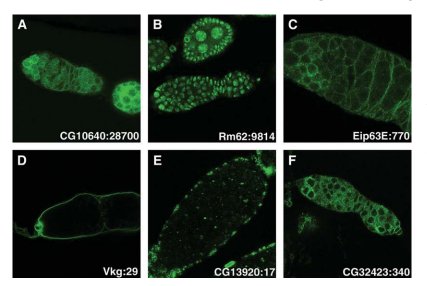


FIGURE 4.—Studies of protein trap expression We compared the apparent intensity of GFPprotein staining in the GSCs with the RNA level of the corresponding gene as determined by Affymetrix arrays (KAI *et al.* 2005). (A–F) The pattern of protein trap expression of the indicated gene (see Table 4 for strain names). The expression level from Affymetrix software (mean of three measurements) is given.

lines whose levels of mRNA are called as "present" on Affymetrix arrays (KAI *et al.* 2005). This suggests that protein trap lines are not limited to a relatively small number of highly expressed genes, but can be used to follow a large fraction of gene activity.

#### DISCUSSION

The Carnegie protein trap collection-a versatile research tool: These experiments significantly expand the number of protein trap lines available for studies of gene expression within a complex multicellular animal (also see accompanying article by QUIÑONES-COELLO et al. 2007, this issue). Our initial characterization of these lines extends previous documentation that the behavior of the EGFP-tagged protein often corresponds to the behavior of the protein to which it is fused. Moreover, we demonstrate that collections such as ours are exceptionally useful as tools of gene discovery. Candidate genes can be selected on the basis of the developmental expression, subcellular localization, or dynamic behavior of particular protein isoforms. The same line can subsequently be used to purify the protein and its associated complexes and to create deletions for further genetic analysis. The Carnegie collection is available for research use from the Carnegie Institution. Information is available at http://www.ciwemb.edu/ resources/proteintrapcollection.html and at http:// flytrap.med.yale.edu/.

**Subcellular distribution of protein location:** Previously, gene and protein trapping in yeast has been used to estimate the fraction of proteins that are localized to various cellular compartments (Ross-MACDONALD *et al.* 1999; KUMAR *et al.* 2002; HUH *et al.* 2003). More than half of all proteins showed a simple localization to the cytoplasm or nucleus. Other subcellular structures labeled by tagged proteins in yeast included the

plasma membrane, the ER, mitochondria, the lysosome, and the perixosome. We obtained similar results. The distribution patterns of EGFP-tagged proteins within Drosophila ovarian cells generally matched those of yeast and most known structures within egg chambers have now been labeled with at least one protein trap line (this study; MORIN *et al.* 2001; CLYNE *et al.* 2003). Moreover, the large size of ovarian cells often allowed us to distinguish the fine structure of several subcellular compartments labeled by EGFP fusion proteins generated in this screen.

Developmental regulation of protein expression: Despite the fact that only one tissue was examined closely, a large number of proteins in the core collection were expressed and many were developmentally regulated. A diverse array of cell types within the germarium including the terminal filament, cap cells, escort cells, germline stem cells, and prefollicle cells were labeled in various lines. However, expression frequently varies from stage to stage, not only in cell type but also in subcellular location, complicating the problem of accurate annotation. Currently, protein trap images within the ovary are being curated in the FlyTrap database. Because of the relative cellular simplicity of the germarium and developing ovarian follicles, it may be possible to develop tools for displaying expression patterns at single-cell resolution in this tissue. It will be particularly valuable to add data from many other tissues and developmental stages for these same lines, to facilitate comparisons.

**Identification of insertions not predicted by genome annotation:** One of the surprising results of our studies was the relatively high frequency of EGFP-positive lines that were located at sites not predicted to fuse to any annotated Drosophila transcript. However, similar results were observed in previous studies of gene trap transposons. At least 44% of insertions analyzed by MORIN *et al.* (2001) were not within annotated genes; moreover, the reading frame of insertions in genes was not determined. In yeast, Ross-MACDONALD *et al.* (1999) observed that while 1346 EGFP-positive insertions were in the correct frame, another 480 were not. Since most lacked an alternative start site, they postulated that a higher than expected frequency of translational frame-shifting may occur. In a recent study in the mouse, 24% of genes were trapped in more than one reading frame (DE-ZOLT *et al.* 2006). We also observed this phenomenon; however, our studies emphasized the difficulty of drawing final conclusions until the location of every insertion and the actual pattern of splicing within the mutant strains have been characterized.

Sensitivity of gene traps: A potential limitation of protein trapping in vivo is that many gene products may be expressed at levels so low that EGFP expressed at the same level could not be detected above background. Only 20% of mouse secretory traps that are G418 resistant express detectable CD2, even though the neophosphotransferase gene is fused to CD2 (DE-ZOLT et al. 2006). Only 33% of  $\beta$ -geo lines resistant to G418 express detectable lacZ. This probably indicates that many genes exist that generate enough neophosphotransferase to confer G418 resistance, but not enough  $\beta$ -galactosidase to be scored as lacZ positive (DE-ZOLT et al. 2006). Consistent with this, Ross-MACDONALD et al. (1999) found that 415 of 1340 in-frame fusions (31%)could be detected above background by immunofluorescence. In contrast, HUH et al. (2003), who tagged complete proteins, detected signals above background for 4156 of 6029 (69%) genes. The system we employed is also designed to tag full-length proteins, and this may have enhanced its sensitivity.

The requirement that each line generate EGFP fluorescence in embryos might provide a limitation on the number of genes that could be tagged. However, our experiments argue that this poses relatively little selection on which genes can be fused. We found that genes expressed in germline stem cells at a wide variety of levels on the basis of microarray studies had been fused in our collection of protein trap lines. There was a rough correlation between the levels observed using antibody staining in these cells and the microarray results. This would indicate that the protein trap methodology can potentially be used to analyze thousands of diverse Drosophila genes.

**Increasing proteome coverage:** Our analysis revealed two major limitations of the current strategy for generating protein traps using P elements. Despite the advantages of embryo sorting, the inherent 5' bias of P-element insertion (BELLEN *et al.* 2004) greatly limits the rate at which new genes can be trapped. Many of the insertions were recovered when an insertion occurred at an internal promoter that lies within a coding intron of another gene isoform. Many genes lack such alternative promoters, so it will be necessary to use different methods to efficiently recover a more diverse collection of protein trap strains.

At least two alternative approaches are worthy of consideration. First, it should be possible to take advantage of the extensive collection of P-element insertions in Drosophila genes that have been generated by the BDGP gene disruption project and other members of the Drosophila community (reviewed in MATTHEWS et al. 2005). We calculate that  $\sim$ 2000 genes already have an existing *P*-element insertion within a coding intron. Moreover, P elements can recombine into the sites of existing Pelements in the presence of transposase (SEPP and AULD 1999). Consequently, a protein trap allele of each of these genes could, in principle, be generated by combining a protein trap insertion of the appropriate reading frame with the "target" gene insertion in a single strain, ideally using inserts bearing scorable markers and then screening for replacement.

Alternatively, transposons with a broader insertional specificity than the P element would be worthwhile. piggyBac elements are suitable for widespread mutagenesis of Drosophila genes (THIBAULT et al. 2004) and as gene traps (BONIN and MANN 2004). Minimal sequences for piggyBac transposition were defined recently (L1 et al. 2005). We obtained several hundred *piggyBac* protein trap insertions that express EGFP, but the *piggyBac* gene trap vector appeared to be less efficient, on the basis of EGFP intensity and Western blotting, than P-element gene traps in nearby locations (also see accompanying article by QUIÑONES-COELLO et al. 2007, this issue). New vectors containing different splice acceptor and donor sites should be tested within the context of a *piggyBac* element. Several new transposons with diverse insertional specificities, including Minos (ARENSBURGER et al. 2005; METAXAKIS et al. 2005), are also worthy of consideration. Continued efforts to provide greater coverage within the Drosophila proteome are warranted because of the exceptional utility of protein traps in analyzing the development and physiology of multicellular organisms.

We thank Lynn Cooley for helpful discussions. We thank X. Morin, W. Chia, A. Hudson, R. Lehmann, and A. Handler for reagents. We are grateful to the following people for their assistance with diverse aspects of the project: Alison Pinder, Joseph Carlson, Martha Evans-Holm, Crista Sewald, Becca Sheng, Melanie Issigonis, Megan Kutzer, Emily Seay, and Dianne Williams. We thank the Howard Hughes Medical Institute, Carnegie Institution, and the National Institutes of Health for support. M.B. was a fellow of the American Cancer Society. T.G.N. is a Howard Hughes Fellow of Life Sciences Research Foundation.

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Communicating editor: T. SCHÜPBACH