Expression of an activated ras gene causes developmental abnormalities in transgenic Drosophila melanogaster

John G. Bishop III and Victor G. Corces
Department of Biology, The Johns Hopkins University, Baltimore, Maryland 21218 USA

We describe the developmental effects of the expression of a normal and mutated Drosophila ras gene ras2 in transgenic Drosophila. A Gly14 → Val14 mutant (Gly14 in ras2 is equivalent to Gly12 in mammalian ras proteins) was constructed in ras2 by site-directed mutagenesis. Inducible constructs of ras2 and ras2Val14 were made by coupling the hsp70 promoter to each of the genes. In addition, the endogenous ras2 promoter was placed upstream of the mutated ras2Val14. These constructs were introduced into the germ line of Drosophila to yield transgenic lines by P-element-mediated transformation. A wide variety of developmental disorders were displayed in flies carrying the mutated ras2 gene driven by either the endogenous or the hsp70 promoter. The basal level of transcription of mutated ras from the uninduced hsp70 promoter was sufficient to produce disturbances in the development of several tissue types. In contrast, phenotypic disturbances were not seen with the normal ras2 gene driven by the hsp70 promoter even when induced by heat shock to very high levels of transcription over normal ras2 levels. A subset of the tissues expressing ras2 during development was particularly sensitive to the expression of ras2Val14. The compound eye was found to develop a dorsal-to-ventral 'scar' correlated with the wave of differentiation occurring in the eye imaginal disc at the time of a single brief induction of the hsp70 promoter ras2Val14 construct.

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The ras proto-oncogenes have fundamental roles in normal and neoplastic cellular processes [reviewed in Barbacid 1987]. Activated ras genes have been found in a substantial percentage of human tumors [Bos et al. 1987; Forrester et al. 1987] and arise experimentally in mouse tumors that have been induced by a variety of treatments such as gamma irradiation [Guerrero et al. 1984]. The ras proteins bind guanine nucleotides, have GTPase activity, and are associated with the plasma membrane [Scolnick et al. 1979; Willingham et al. 1980; Gibbs et al. 1984; McGrath et al. 1984; Sweet et al. 1984; Willumsen et al. 1984; Manne et al. 1985; Temeles et al. 1985]. Most amino acid substitutions in the ras protein at position 12 result in a lowered rate of GTP hydrolysis and a transforming (or activated) phenotype [Willingham et al. 1980; Gibbs et al. 1984; McGrath et al. 1984; Seeburg et al. 1984; Sweet et al. 1984; Willumsen et al. 1984; Chipperfield et al. 1985; Manne et al. 1985; Temeles et al. 1985]. Sequence comparisons between ras and G proteins suggest that ras proteins act in transducing signals across the cellular membrane [Hurley et al. 1984; Lochrie et al. 1985; Tanabe et al. 1985]. It has been found that ras proteins injected into mammalian cells cause a change in morphology, membrane ruffling, and increased pinocytosis [Bar-Sagi and Feramisco 1986].

Evolutionarily conserved ras genes have been found in yeast, slime mold, fish, mammals, plants, and Drosophila [Barbacid 1987]. In yeast, gene disruption experiments have shown that ras proteins are involved in the regulation of growth and the activation of adenylyl cyclase [Kataoka et al. 1984]. The ras genes are expressed in both proliferating and terminally differentiated cells in Drosophila [Segal and Shilo 1986] and in humans [Chesa et al. 1987]. Dictyostelium transformed with activated ras genes showed abnormal aggregation; wild-type ras genes had no effect [Reymond et al. 1986]. In mammals, ras involvement in differentiation appears to be complex because injection of ras oncogenic protein has triggered morphological differentiation [neurite outgrowth] of PC12 cells [Bar-Sagi and Feramisco 1985], whereas the differentiation of a skeletal myoblast cell line can be prevented with the oncogenic forms of H-ras and N-ras genes [Olsen et al. 1987]. Transgenic mice carrying a MMTV LTR-promoted-v-ras construct showed in developmental aberration of a specific tissue, the Harderian gland, and tumors after varying latency periods [Sinn et al. 1987]. The choice of promoter in these studies was of great importance because transgenic mice carrying the promoter for pancreatic elastase driving an activated ras gene developed pancreatic tumors with no latency [Quaife et al. 1987] while transgenic mice carrying the whey acidic protein gene promoter driving ras have a very low incidence of tumors [Andres et al. 1987].

We were interested in several questions involving ras genes and Drosophila. First, what phenotype[s] would
evolve when a mutated form of ras is incorporated into the genome of an insect; would tumors or abnormal growth result? Second, if ras affects the normal development of a whole organism, would all tissues be affected in a similar manner or degree; would the sensitivity be time dependent? Third, what would be the consequence of an increased expression of the normal ras gene?

Results

Structure of the Drosophila ras2 locus

At the outset of these experiments, crucial information necessary for the construction of functional transforming ras genes was lacking. Partial genomic DNA sequences were available (Neuman-Silberberg et al. 1984; Mozer et al. 1985) and a preliminary Northern blot analysis indicated that three transcripts of 2.6, 2.1, and 1.6 kb originated from the ras region (Mozer et al. 1985). Did all three transcripts originate from the ras2 gene or were they unrelated? As exogenous promoters were to be used to drive ras2 transcription, the start of transcription of the ras2 gene had to be defined more precisely. Because several potential polyadenylation sites had been recognized during the course of the genomic DNA sequencing, the determination of the processing site(s) actually used was essential for successful transformation experiments.

A detailed Northern blot analysis established that the larger transcripts are different from the ras2 gene and map to the 5' side of ras2. The analysis and sequencing of ras2 cDNA clones revealed the probable transcription start site and the polyadenylation site of the ras2 gene. The structure of the ras2 gene obtained from genomic and cDNA sequencing, and Northern blot analysis is summarized in Figure 1. The 0.86 kb Clal fragment [Fig. 1B], mapping 55 nucleotides upstream of the putative ras2 transcription start site (detailed below and Fig. 2), and extending in the 5' direction from ras2 sequences, hybridized only to the 2.6 and 2.1 kb RNA species on Northern blots [S. Parkhurst and V. Corces, unpubl.]. On the other hand, the HindIII–BamHI fragment, comprising the bulk of the ras2 coding region, contained in pUC8-HB-1.2 [Fig. 1B], hybridized to a single 1.7-kb RNA species. Segal and Shilo (1986) identified a 1.6-kb transcript using a similar probe. From these results, we conclude that the two large transcripts map close to the 5' end of the ras2 gene but are different from ras2.

The two ras2 cDNA clones depicted in Figure 1C revealed a potential transcription initiation site. The independent clones had identical 5' terminal sequences and ended at the position marked 1 in Figure 2. A potential TATA sequence, GATATT, was found at position −55 in the genomic sequence. O'Connell and Rosbash (1984) found the same sequence at −51 from the start of transcription of the Drosophila ribosomal protein 49 gene. In

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**Figure 1.** Scale drawings of the ras2 genomic organization, subcloned fragments, and cDNA clones. (A) A genomic restriction map of the ras2 locus was determined from several subcloned genomic fragments. The arrow shows the direction of transcription of ras2. (B) Subcloned fragments used in this study. (C) Two cDNA clones rasCDNA2-1 and rasCDNA2-2 of the ras2 gene were isolated. Open reading frames were found in the sequence upstream of ras2 in the opposite transcriptional orientation. No strong homologies were found with the conceptual polypeptides encoded therein and the BIONET protein database (Release 14).
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A

| -160 | -150 | -140 | -130 | -120 | -110 |
|------|------|------|------|------|------|
| AGAATTTATCTATTTCTGGCCAATT TTCTGTGTTTTTCCTGATGACCTGCTT |
| -100 | -90  | -80  | -70  | -60  | -50  |
| TAGTGATGGGCGGAGCTAGTGGCTACGGATAGTTGATAGCTGATATTCTTTT |
| -40  | -30  | -20  | -10  | 10   | 20   |
| CAGAGACCGCGAACCGGTCCTGCCCAGGCTGGATTTTCTCAGTTAACCGAGAC |
| 30   | 40   | 50   | 60   | 70   | 80   |
| GCGCTGCTCGGAGGCAAGAAAATAAAAATATTTTCTGGCCAACCTGTCAGTTTTA |
| 90   | 100  | 110  | 120  | 130  |
| GGGGGGAAATTCGATGGCAACTACATGCTGCTGGCCGCGGCGGGCGGGCGGAG |
| MetGlnMetGlnThrTyrLysLeuValValGlyGlyGlyGlyValGlyLys |

B

| 870  | 880  | 890  | 900  | 910  | 920  |
|------|------|------|------|------|------|
| AACGATGGAACATCGGATGYYGAGGCGAAATGCAGAGGAGTAAGCTCTTTTTTCA |
| 930  | 940  | 950  | 960  | 970  | 979  |
| cttaccagattcatataacctgatatatttaacctagATTGGACACGGGCAGCAC |

Figure 2. (A) Genomic DNA sequence of the promoter region of ras2. The putative CAAT sequence is boxed at position -141 to -133 and the putative TATA sequence is boxed at position -55 to -50 (Corden et al. 1980). The first 18 codons of ras2 are shown starting at position 87, the conceptual translated amino acid sequence is shown below the line. (B) Genomic DNA sequence and predicted amino acid sequence of the exon 2-intron-exon 3 region of ras2. Numbering of positions was based on the cDNA clones 5' termini; this is the putative mRNA start site. A putative second site at -29 is also underlined. Restriction sites are shown by heavy underlining. The cDNA sequence of the two ras cDNA clones revealed the splicing model of Mozer et al. (1985), based solely on genomic sequencing, required modification. The corrected splicing scheme and revised conceptual polypeptide sequence are shown. Introns sequences are shown in lower case.

addition, a eukaryotic CAAT sequence was discovered in the sequences immediately upstream of the ras2 gene [Fig. 2].

One of the cDNAs was full length and allowed the identification of the processing site for polyadenylation. Clone rascDNA2-1 [Fig. 1C] ended in a polyadenylate sequence 36 nucleotides downstream from a polyadenylation signal sequence AATAAA [Proudfoot and Brownlee 1976]. This was the most distal of the potential sites identified in the genomic sequencing; the other two lay 45 and 683 nucleotides closer to the ras2 coding sequences. Functional transformation with ras2 gene constructs would most likely require this polyadenylation signal and flanking sequences for proper processing. A ras2 transcript of 1614 nucleotides, not including the polyadenylate tract, is predicted from the cDNA and genetic analysis, and was in close agreement with experimental results.

To obtain additional 5' and 3' sequences for the transformations with the ras2 gene, a 4.6-kb Xhol ras2 fragment was isolated from a Canton S library [Maniatis et al. 1978] and subcloned as pBl20-4.6 Xhol [see Fig. 1B]. The fragment contains 2 kb of sequence extending 5' of the initiation codon and 400 nucleotides in the 3' direction beyond the polyadenylation site.

**ras2 gene mutagenesis and P-element constructs**

The three constructs made for the present study are shown in Figure 3. The hsp70 promoter was coupled to the ras2 4.6-kb Xhol gene fragment at the Hpal site located 83 nucleotides upstream of the initiation codon of ras2 gene and cloned into the P-element vector Carnegie20 [C20] (Rubin and Spradling 1983) to create C20-29 [Fig. 3A]. Oligonucleotide-directed mutagenesis (Zoller and Smith 1983) was employed to create the mutation of Gly14→Val14 (Gly14 in ras2 is equivalent to Gly12 in mammalian ras proteins; Mozer et al. 1985) to create ras2Val14, this 4.6-kb Xhol fragment mutated at a single nucleotide was subcloned into C20 and named C20-35 [Fig. 3B]. The ras2Val14 gene construct was coupled to the hsp70 promoter at the Hpal site located 83 nucleotides upstream of the initiation codon of the ras2Val14 gene to create C20-42 [Fig. 3C].

The C20 constructs with helper plasmid pnr25.7 supplying transposase in trans were injected into rosyD06 embryos with a Canton S background. These flies were mated to the rosyD06 parental stock and their progeny scored for the presence of wild-type eye pigmentation indicating transformation with C20 which has a wild-type rosy gene. This protocol typically results in a single integrated copy of the gene of interest [Rubin and Spradling 1983]. At least three separate lines were developed for each construct [see Table 1]; each of these lines were crossed to chromosome balancer stocks with the dominant markers Curly on the second, Bar on the X, or Stubble on the third chromosome to map the chromosomal location of the integrated transposon. Crosses to obtain homozygous stocks were done where possible, these results are shown in Table 1.
Figure 3. *ras2* gene constructs used for germ line transformations. In each case, the *ras* gene was inserted into the *Sall* site of Carnegie20 (C20) in an opposite transcriptional orientation to the *rosy* gene (A) C20-29. The *ras2* gene was coupled to the hsp70 promoter at a *Hpal* site located 83 nucleotides upstream of the initiation codon. (B) C20-35. The first exon of *ras2* was subjected to oligonucleotide-directed mutagenesis (Zoller and Smith 1983) to create the mutation of Gly14 — Val14 and installed in pIB20-Val14-4.6. The single base change is indicated as a large letter. The *Xhol* fragment was then subcloned into C20 to yield C20-35. (C) C20-42. The hsp70 promoter was installed into pIB20-Val14-4.6 *Xhol* after the normal promoter was removed. The 3-kb fusion *Xhol* fragment in pIB20-hsp70-*ras2*Val14 was then subcloned into C20.

Heat-shock promoted *ras2* constructs C20-29 and C20-42 in transgenic *Drosophila* were inducible by heat shock

The inducibility of the hsp70 promoter fusion constructs was measured by Northern blotting of RNA isolated from adults of the C20-29 and C20-42 transgenic lines before and after heat shock and probing with *ras* subclone pUC8-HB-1.2 kb. Total RNA was isolated from adults that were kept at 22.5°C or were heat shocked for one hr at 37.0°C. The autoradiograph is shown in Figure 4. In the RNA samples from the heat shocked (shown as HS in Fig. 4) animals carrying hsp70-promoted *ras* genes, inducible transcripts were detected. Because rRNA co-migrates with the major *ras* transcript and hinders transfer we could not estimate the actual degree of induction. The origin of the larger, inducible RNA transcript is unknown.

An S1 nuclease assay was developed to exploit the sequence differences between the endogenous *ras2* transcripts and the heat shock-fusion transcripts. The hsp70 promoter was expected to initiate transcripts 223 nucleotides upstream of the *Hpal* site where the hsp70 promoter was fused to the *ras2*Val14 sequence in the C20-42 constructs as shown in Figure 5A. Endogenous *ras2* transcription was known to initiate upstream of this *Hpal* site from sequence analysis of the *rascDNAl* clones. The 559-bp *Xhol*-Aahall fragment from pIB20-*ras2*Val14-Val70 was end-labeled at the *Aahall* site on the anti-sense strand. The cleavage site for *Aahall* was 122 nucleotides downstream from the *Hpal* junction site so that 122 bp protected fragments would be expected from endogenous *ras2* RNA and 345 bp fragments would be expected from hsp70-promoted transcripts (see Fig. 5A). Line C20-42PG1-4X was assessed for its ability to produce hsp70 promoted transcripts upon heat induction by S1 nuclease protection experiments and the results are shown in Figure 5B. High levels of transcripts were observed after heat induction from the hsp70-*ras2*Val14 con-
Total RNA was isolated from hspZO-promoted fragments from pUC8-HB-1.2 kb representing most of the third exon of rRNA co-migrated with rRNA, lasl. The lower transcript and interfered with the transfer.

The hsp70-ras2Val14 construct caused development disturbances in both induced and uninduced animals; the hsp70-ras2 construct caused no observable disturbances.

The ras2Val14 gene driven by the hsp70 promoter was introduced into three transgenic lines of the C20-42 series (see Table 1). Initial observations indicated each of these stocks had low fertility, laying many eggs that failed to hatch, and showed developmental disturbances in their wing and eye structures even when kept at 18°C or 22.5°C throughout their entire development. Bristle formation was also affected in that the bristles were of variable size and placement. This was particularly noticeable on the wing anterior margin. Sternopleural bristles, and humeral bristles of the thorax were often bent or forked. The cuticle region of the humerus was also pronounced in many of the flies. The basal level of transcription from the heat shock promoter seen in the S1 nuclease protection assay (see Fig. 5B) was presumably responsible. The fly compound eye was a particularly sensitive gauge because its orderly, almost crystalline, nature readily revealed subtle defects to the observer. Conversely, the C20-29 series of lines carrying the hsp70-promoted normal ras2 gene appeared entirely unaffected when reared under the same temperature regimes.

In the next series of experiments, animals from each line bearing the hsp70-ras2Val14 heat-inducible construct were studied for their sensitivity to high-levels of transcripts induced at defined and brief periods during their development. Animals were mass cultured and heat shocked a single time at 37.0°C for 1 or 2 hr unless specified otherwise. This amount of time represents 0.5–1% of the generation time of D. melanogaster under these conditions. Adult flies, pupae, and larvae from each line and from the rosy-506 control line were scored for visible phenotypes appearing daily over at least one generation time period. The wings of animals heat shocked during the third instar and pupal stages failed to fuse at the dorsal and ventral surfaces resulting in a bloated appearance to the wings. Wings were typically smaller and distorted; they were absent and thoraces were distored in a low percentage of flies although usually on one side of the fly. The same phenotype resulted from a pulse of as short as 5 min at 37.0°C; even heat pulses of 1 hr at 35.0°C caused the unfused wing phenotype. The sensitive period for wing anomalies was found to be quite broad and spanned the early third instar to mid-pupal stages. Yet animals heat shocked earlier in their development rarely had wing defects.

The hsp70-ras2Val14 animals taken from the entire third instar stage of development and subjected to a 2-hr heat shock had an overall survivability of 20–40%; the stage at which death occurred was highly variable ranging from within hours resulting in blackened larval carcasses, to 6 to 7 days later near or at adult emergence. However, almost all animals heat shocked in the narrower window of time from a few hours prior to the onset of pupariation and for a few hours thereafter never survived. These animals died as larvae and pupae with missing abdominal structures to incompletely everting head tissue, grossly aberrant wings, and distorted legs.

Strikingly, in these animals bearing the hsp70-ras2Val14...
construct, dorsal-to-ventral 'scars' developed in the middle-to-posterior portion of the adult eye when the animals were heat pulsed from 1 to 2 days before the onset of pupariation (late-third instar stage). Nearly all animals that developed to pharate adults or actually eclosed developed these scars bilaterally. Scars were observed in the late pupal stages as unpigmented regions of the eye visible through the pupal case or upon dissection. The scars consisted of unpigmented, irregular, and fused ommatidia as shown in Figure 6C and D. The adult shown in Figure 6D was heat shocked from 22 to 24 hr before pupariation, animals heat shocked earlier in the third instar stage developed scars more posterior to those shown in Figure 6C and D. Control adults of the parental stock ry506 never displayed any phenotypic effects from heat shock pulses of 2 hr, during any portion of their development, when incubations were kept exactly at 37.0°C. Heat pulses of 1 hr to animals of the C20-42 line bearing the hsp70-ras2 constructs caused less severe scarring (see Fig. 6B) and pulses of less than 30 min failed to produce scars. Using the optical neutralization technique, observations of adult eye photoreceptor patterns in animals heat shocked during development and bearing roughened eyes indicate that often the ommatidia were fused resulting in ommatidia of twice normal size. Where photoreceptors could be identified, two juxtaposed sets of photoreceptors were found and each set was composed of the proper number (data not shown). The technique is limited and does not detect absent photoreceptor clusters. This would indicate that the pigment cells in these fused ommatidia failed to develop or, move properly, to optically insulate each photoreceptor cluster resulting in two bundles of photoreceptors in one large ommatidium. The single lens that formed presumably results because the cone cells secreting the lens were able to compensate.

Somewhat surprisingly, high transient levels of normal ras2 transcripts were tolerated well in the heat-shocked lines bearing hsp70-ras2 constructs. No visible phenotypic resulted and 98–100% of the animals subjected to a single 2 hr heat survived with no discernible ill effects. The same result was observed in animals of the control line ry506.

The ras2Val14 gene driven by the endogenous ras2 promoter caused developmental disturbances similar to those seen with the hsp70 promoter

The developmental growth disturbances seen in the hsp70-ras2Val14 animals may represent the accumulation of ras2Val14 transcripts in tissues that never express ras2 since the hsp70 promoter is active in all or almost all the tissues of the fly, whereas ras2 was seen to be expressed in a temporal and tissue-specific manner (Segal and Shilo 1986). We could test if the animals were sensitive to low levels of ras2Val14 by utilizing a construct with 2 kb of upstream sequences from the endogenous ras2 gene. In this way, low levels of ras2Val14 product would be produced in presumably the same tissues as the normal ras2 product. These animals could then be compared to the uninduced hsp70-ras2Val14 flies to see if any of the growth disturbances were shared or dissimilar. We assumed that the controlling sequences for proper levels of ras2 transcription map quite close to the gene since (as shown above) different unrelated transcripts were encoded within 1 kb of the upstream regions from ras2. The inclusion of 2 kb of upstream sequences would most probably allow proper transcription of the ras2 construct. Since we had previously seen that even high levels of the normal ras2 transcripts induced by heat shocks had failed to yield any phenotypes, we presumed that low levels being driven from the endogenous ras2 promoter would have no effect on growth or fertility. No phenotype had been associated with lines bearing the hsp70-promoted normal ras2 construct whether induced
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Figure 6. Phenotypes of the ras2val14 transformants. Anterior is to the left and dorsal is at the top. [A] Scanning electron micrograph (SEM) of a wild-type adult Drosophila eye. [B] SEM of an adult eye from line 42PG1-4X heat shocked 1 hr at 37.0°C during the third instar developmental stage. (C) SEM of an adult from line 42PG1-4X heat-shocked for 2 hr at 37.0°C during third instar stage. [D] Light micrograph of adult from line 42JGB3-A heat shocked 2 hr at 37.0°C during the third instar stage 22 to 24 hr before pupa formation.
with heat shocks or from the basal level of transcription from the hsp70 promoter.

The eight lines carrying the construct C20-35, i.e., the 2 kb of endogenous sequences upstream of ras2 driving a mutated ras2Val14 gene, did display developmental anomalies, albeit to different degrees. All lines showed reduced viability and fertility, especially when homozygous, of a similar nature to the hsp70-promoted ras2Val14 construct bearing lines of the C20-42 series. The gene-dosage effect was also seen in trans-heterozygotes derived from crosses between lines C20-35-7 and C20-35-14 as well as lines C20-35-7 and C20-35-3, indicating that the reduced viability and poor development resulted from the levels of ras2Val14 and not solely the position of insertion. Typically, the wings of most of the flies developed poorly. They were held outstretched and the ventral and dorsal surfaces of the wing failed to fuse, often blackening as the flies aged. Ommatidia of the eyes were slightly disordered making the eyes rough in appearance in the majority of the individuals of the most severely affected lines; again, this was similar to the phenotype observed in the most affected flies from the uninduced hsp70-promoted ras2Val14 lines. Lines C20-35-7 and C20-35-14 were weak, slow in movement, and showed a complete loss of wing beating whereas individuals of line C20-35-20 retained a diminished ability suggesting the construct was susceptible to position effects. This poor wing movement phenotype was more severe than the similar phenotype seen in the animals from the noninduced hsp70-promoted ras2Val14 lines. In addition, the bristle formation on the wing and thorax, and the protruding cuticle growth of the humeral region observed in lines of the C20-35 series resembled phenotypes observed in the uninduced C20-42 series bearing the hsp70-ras2Val14 construct.

Discussion

Transient high levels of normal ras2 transcripts produced in the heat-inducible C20-29 lines, or the low basal levels produced in uninduced flies, do not seem to have any deleterious effects on the flies.

On the other hand, high levels of ras2Val14 transcripts induced in animals carrying the heat shock-inducible constructs (C20-42 series) caused a wide range of developmental anomalies. The low levels expressed basally from the uninduced hsp70 promoter driving ras2Val14 also caused developmental anomalies of a much lesser degree, which appeared to be quite similar to the aberrant phenotypes seen in animals carrying the normal ras2 promoter driving ras2Val14. The heat shock-inducible phenotypes may represent the primary effects of the expression of the ras2Val14 gene in the visibly affected tissues, although the phenotypes could be caused by transcription in tissues which then influences the tissues we see disrupted. When the hsp70-promoted ras2Val14 construct lines were reared at 18°C or 22.5°C, they did show low fertility, poor wing development, and subtle defects in the eye. Also, bristle malformation was particularly noticeable on the wing anterior margin.

cause we see similar phenotypes in animals with the endogenous promoter, i.e., the 2 kb of upstream sequences from the endogenous ras2 locus, we feel that the phenotypes result from the sensitivity of those tissues normally expressing ras transcripts, although we have no proof to demonstrate that the 2 kb of upstream sequences are sufficient for proper tissue and temporal regulation of ras transcripts. Ostensibly, the ras2 promoter in the constructs might be truncated or altered in some way that resulted in a release from the normal transcriptional regulation of ras2 causing it to mimic the hsp70 promoter. A more likely consequence of such a truncation would be the down-regulation of transcription resulting in the absence of any discernible phenotype. Since phenotypes were observed the promoter must be active and is probably functioning in a normal manner.

We have chosen to focus on the eye phenotype because the development of the Drosophila eye has been well studied both genetically and ultrastructurally, and the eye consists of a relatively small number of different cell types. During the third instar and early pupal stages, a wave of photoreceptor cell differentiation visible as a morphogenetic furrow, moves across the eye imaginal disc from the posterior to the anterior margin [Ready et al. 1976]. Clusters of photoreceptor cells form initially, later cone, pigment, and hair nerve cells are added sequentially to form an ommatidium made up of 22 cells [Ready et al. 1976; Tomlinson and Ready 1987]. Genetic analysis of clones and observations of eye development at the cellular level indicate that the cells forming the ommatidia differentiate using positional cues to guide their decisions [Ready et al. 1976; Lawrence and Green 1979; Tomlinson and Ready 1987]. To explain the disordered development of the eye that results from the high levels of ras2Val14 in light of the prevailing models for ras function, we might presume that the phenotype arises because of the aberrant transduction of a signal across the membrane of the eye cells. This signal serves to cue the cells to adopt their appropriate morphology. A considerable number of models are also likely. The dorsal-to-ventral scar could result from a disruption in the transduction of extracellular signals by the photoreceptor cells as they begin to differentiate, arresting or altering their development. Alternatively, the photoreceptors may fail to elaborate the proper positional cues needed by neighboring cells of the pigment or cone cell types that undergo subsequent differentiation [Tomlinson and Ready 1987]. Analysis of the third instar larval eye disc for photoreceptor patterning, as well as the pupal eye discs, when the pigment and cone cells are moving into position and differentiating, from hsp70-ras2Val14 animals that are heat shocked in the larval stage may allow us to identify the particular cell types that appear sensitive to ras levels. The sensitivity of the eye to ras2Val14 expression is interesting in view of recent work indicating signal transduction via a membrane-associated tyrosine kinase molecule as a cue for photoreceptor differentiation. Sevenless, as eye-specific gene [Harris et al. 1976], when mutant, causes the seventh photoreceptor neuron to become homeotically trans...
formed into a nonneuronal cone cell. *Sevenless* encodes a putative tyrosine kinase present at the cell membrane [Banerjee et al. 1987; Hafen et al. 1987]. Perhaps the product of ras2 is a downstream molecule involved in this or other transduction events occurring during the differentiation of the eye.

Two genes have been identified that influence the adult phenotype of the eye during the same development time period and cause a dorsal-to-ventral scar; X-rays can cause similar scars [Becker 1957]. An allele of Notch [Foster and Suzuki 1970], which encodes a membrane protein with homology to epidermal growth factor [Wharton et al. 1985; Kidd et al. 1986], and an allele of Shibere [Poodry et al. 1973], thought to be deficient in membrane recycling [Kim and Wu 1987], develop a dorsal-to-ventral scar of the adult eye if the third instar larvae when reared at the nonpermissive temperature for a period of several hours. A dorsal-to-ventral scar that is lacking pigmentation (see Fig. 6D) as seen with the ras2val14 mutation has not, to our knowledge, been described before and suggests to us that dorsal-to-ventral scars may be formed in a variety of ways. Henkemeyer et al. [1987] have found that the Abelson proto-oncogene [abl] of Drosophila, when mutant, causes the retinal pattern of the adult eye to be disrupted and retinal cells are often missing and ommatidia can fuse. Experiments are now in progress to investigate possible interaction among ras2val14, Notch, and sevenless alleles.

The particular pattern of tissue sensitivity of ras2val14 expression in Drosophila is correlated to the presence of high levels of endogenous ras expression in the particular organs of the developing fly [Segal and Shilo 1986] but is limited to a subset of these tissues. In particular, the tissues in the larval stages that are proliferating, the imaginal discs, gonads, and brain, express each of the ras genes. In the adult, the ovaries and the terminally differentiated cortex of the brain and ganglia express ras transcripts [Segal and Shilo 1986]. Our results indicate that the eye and wing primordia are particularly sensitive to ras2val14 expression. The low fertility of the females may reflect the expression of mutant ras2 in the ovaries which normally produce high levels of ras transcripts (Segal and Shilo 1986). In Drosophila, as also shown in the mouse [Sinn et al. 1987], there appear to be specific tissues in which an activated ras gene is mute and others in which gross developmental anomalies occur. This may result from the fact that the different forms of ras play different roles in the separate tissues; alternatively, ras proteins may play unequal roles in the development of each of the tissues [Sinn et al. 1987].

It will be interesting to see to what degree other proto-oncogenes that have been described in Drosophila [reviewed in Shilo 1987] will affect development when mutated and transformed into the germ line of Drosophila. The lines developed in this study could then be used to study possible synergistic interactions among oncogenes as has been done in mice [Sinn et al. 1987]. The nature of the molecules that interact with ras proteins and the mechanisms of their actions are at present unknown in multicellular organisms. These mutant lines will be the starting material for a genetic screen to isolate revertants in the ras2 phenotypes caused by second-site mutations; these mutations should identify genes involved with ras functioning.

Materials and methods

Preparation of nucleic acids, genomic and cDNA library screening, and DNA sequencing

Plasmid DNA isolations, ligations, and gene library screens were done by standard methods [Maniatis et al. 1982]. Total RNA was extracted from adult Drosophila using the SDS-phenol technique of Spradling and Mahowald [1979]. The cDNA clones were isolated from the Kauvar library [Poole et al. 1985]. Denaturing formaldehyde gels were used for RNA gels [Maniatis et al. 1982] and RNA transfers were to nylon membranes [ICN Biotrans]. DNA sequencing and end-labeling were done by the methods of Maxam and Gilbert [1980].

Constructs and oligonucleotide-directed mutagenesis

C20-29: An XhoI–BanHI fragment of 437 bp from the Drosophila hsp70 promoter containing sequences from −194 to +233 was isolated from pVC8-1 (V. Corces, unpubl.), filled in at the BamHI site and used to replace the S′ XhoI–Hpal fragment containing the normal ras2 promoter in plBI20-4.6Xhol to create plBI20-hsp70-ras2. Its 3.0 kb XhoI fragment was subcloned into Carnegie20 to make C20-29.

C20-35: The Hpal–HindIII fragment from plBI20-4.6Xhol containing the first and second exons of ras2 was subcloned into plBI20. Oligonucleotide-directed mutagenesis was performed as described (Zoller and Smith 1983). The single base change creating the mutation at codon 14 of ras2 is underlined in the oligonucleotide sequence CCGGGTGCGGCTGGGCA. The oligonucleotide spans positions 121–138 in Figure 2 and was annealed to the single-stranded DNA from plBI20-Hpal–HindIII. Screening of mutant clones was done with the end-labeled oligonucleotide on DNA isolated from single colonies and spotted onto nitrocellulose. One putative clone was sequenced to verify the presence of the mutation which creates an AhalI site. The mutated Hpal–HindIII fragment was inserted in place of the wild-type fragment to create plBI20-Vfli4-4.6Xhol and the XhoI fragment was inserted into C20.

C20-42: The XhoI–BamHI hsp70 promoter was installed into plBI20-Vfli4-4.6Xhol after the 2-κb S′ XhoI to Hpal fragment was removed by complete digestion with Hpal followed by partial digestion with XhoI. The 3-κb fusion was then subcloned into C20.

S1 nuclease protection assay

The 559 bp XhoI–AhalI fragment from plBI20-hsp70-ras2val14 was end-labeled at the AhalI site on the anti-sense strand. End-labeled probe (50 ng) was hybridized overnight at 46°C in S1 hybridization buffer [Maniatis et al. 1982] to 50 μg [with the exception of lane G which contained 150 μg] of total RNA isolated from adult flies that had been reared at 22.5°C (controls) or kept at 37.0°C for 1 hr (heat shocked). After S1 nuclease digestion of the hybrids, the nucleic acids were recovered by ethanol precipitation and loaded on a 5% denaturing acrylamide gel [Maniatis et al. 1982]. Unless otherwise indicated, all heat shocks were performed at 37.0°C by immersing vials containing flies of varying stages in a Lauda water bath.
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J G Bishop and V G Corces

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