Raf acts downstream of the EGF receptor to determine dorsoventral polarity during *Drosophila* oogenesis

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In *Drosophila*, as in mammalian cells, the Raf serine/threonine kinase appears to act as a common transducer of signals from several different receptor tyrosine kinases. We describe a new role for Raf in *Drosophila* development, showing that Raf acts in the somatic follicle cells to specify the dorsoventral polarity of the egg. Targeted expression of activated Raf (Raf\(^{act}\)) within follicle cells is sufficient to dorsalize both the eggshell and the embryo, whereas reduced Raf activity ventralizes the eggshell. We show that Raf functions downstream of the EGF receptor to instruct the dorsal follicle cell fate. In this assay, human and *Drosophila* Raf\(^{act}\) are functionally similar, in that either can induce ventral follicle cells to assume a dorsal fate.

[Key Words: Raf serine; threonine kinase; oogenesis; dorsoventral signaling]

Received November 11, 1993; revised version accepted December 23, 1993.

The *Drosophila* ovary is made up of ~16 ovarioles, each consisting of a series of egg chambers [King 1970; Mahowald and Kambysellis 1980], that have been subdivided into 14 distinct stages [King 1970]. Until mid-oogenesis, the egg chamber is fairly symmetrical with respect to the dorsoventral position of the oocyte nucleus and to the shape of the overlying follicle cells. During stage 8/9, the oocyte nucleus migrates to the prospective dorsal side of the oocyte, and subsequently, the dorsal follicle cells become more columnar and tightly packed. The follicle cells eventually secrete the eggshell or chorion, which is itself asymmetric as indicated most clearly by the presence on the anterodorsal side of the dorsal appendages.

The dorsoventral polarity of the *Drosophila* egg is established by means of signaling between the germ line-derived oocyte and the surrounding somatic follicle cells [Schupbach 1987; Schupbach et al. 1991]. The current model [e.g., see Schupbach et al. 1991] postulates that an asymmetric signal, from the oocyte, instructs the follicle cells nearest the oocyte nucleus to assume a dorsal fate. The follicle cells, in turn, establish the dorsoventral polarity of the developing embryo by restricting the activation of a ligand to the ventral side of the embryo, where it specifies embryonic dorsoventral polarity [Schupbach 1987; Manseau and Schupbach 1989; Stein et al. 1991; Stein and Nusslein-Volhard 1992].

Of the maternal effect genes involved in dorsoventral signal transduction, *gurken* is most likely to encode the dorsal signal [Schupbach 1987; Neuman-Silberberg and Schupbach 1993]: the *gurken* transcript is localized asymmetrically, concentrating in the anterodorsal region of the oocyte. Furthermore, *gurken* has been shown to encode a TGF-α homolog, lending further support to its potential role as a ligand. *fs(1)K10*, *spire*, *cappuccino*, and *squid* [Wieschaus et al. 1978; Prost et al. 1988; Manseau and Schupbach 1989; Kelley 1993] act to confine the signal to the dorsal side of the oocyte [Neuman-Silberberg and Schupbach 1993]. The *Drosophila* epidermal growth factor (EGF) receptor homolog, encoded by *torpedo* or *DER* [Price et al. 1989; Scheijer and Shilo 1989], is required in the follicle cells to receive the signal from the oocyte [Schupbach 1987; Clifford and Schupbach 1989; Schupbach et al. 1991]. In addition, a putative transmembrane protein, encoded by the gene *rhomboid*, may potentiate the interaction between the EGF receptor and its ligand [Ruohola-Baker et al. 1993].

*DER* not only specifies the dorsal follicle cell fate, but also determines the dorsoventral axis of the developing embryo. On the ventral side of the egg chamber, a signaling cascade is initiated that culminates in the localized activation of the *Toll* receptor in the embryo after fertilization [Anderson et al. 1985; Hashimoto et al. 1988, 1991; Stein et al. 1991; Chasan et al. 1992; Stein and Nusslein-Volhard 1992]. Activating *DER* within the dorsal follicle cells may block production of an active ligand for *Toll*, and in so doing specify the dorsal side of the embryo.

Here we demonstrate a role for the Raf serine/threonine kinase in establishing dorsoventral polarity during oogenesis.
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oogenesis. Raf has been implicated in at least two different signal transduction pathways in *Drosophila*: (1) in relaying signals from the *torso* receptor tyrosine kinase (RTK) to determine cell fates at the embryonic termini [Ambrosio et al. 1989a,b] and (2) in signaling from the *sevenless* RTK to specify the fate of the R7 photoreceptor cell in the developing eye [Dickson et al. 1992]. Activating *torso* or *sevenless* initiates a signaling cascade that results in the activation of Ras1, possibly by repressing Gap1 (a Ras GAP) and stimulating Sos [a guanine nucleotide exchange factor] [Rogge et al. 1991; Simon et al. 1991; Dickson et al. 1992; Fortini et al. 1992; Gaul et al. 1992; Lu et al. 1993; Doyle and Bishop 1993]. Ras in turn activates Raf that, in the terminal pathway, appears to activate the mitogen-associated protein (MAP) kinase activator, MAP kinase kinase, or MAPK/ERK Kinase (MEK) [Tsuda et al. 1993].

Using targeted expression, we show that Raf functions downstream of the EGF receptor to transduce the germ-line signal that specifies the dorsal follicle cell fate and establishes the dorsoventral polarity of the developing embryo. We have expressed activated Raf (Raf<sup>act</sup>) within follicle cells, and demonstrate that ectopic activation is sufficient to dorsalize both the egg shell and the embryo. Conversely, reduced Raf activity is shown to ventralize the eggshell. We demonstrate that human and *Drosophila* Raf<sup>act</sup> are functionally similar, in that either can induce ventral follicle cells to assume a dorsal fate.

**Results**

*Reducing Raf activity leads to ventralized eggs*

We investigated whether Raf acts in signal transduction during oogenesis, in particular in the EGF signaling pathway. In *Drosophila*, Raf has been shown to act downstream of at least two RTKs, *torso* and *sevenless* [Ambrosio et al. 1989b; Dickson et al. 1992], but as yet not the EGF RTK. If Raf is required in dorsoventral signaling, then lowering the level of expression of Raf, or its activity, should lead to ventralized eggs resembling those produced by EGF receptor mutant females.

To obtain adult females with which to investigate possible oogenesis defects, we used a hypomorphic *Draf* mutation (*Draf<sup>HMT</sup>*; see Materials and methods) [Kramers et al. 1983; Melnick et al. 1993]. Whereas many *Draf* mutations behave as nulls and result in larval–pupal lethality, *Draf<sup>HMT</sup>* mutants survive to adulthood and are fertile (Melnick et al. 1993; Perrimon et al. 1985). *Draf<sup>HMT</sup>* mutants exhibit a rough eye phenotype consistent with the role of Raf in the *sevenless* RTK signaling pathway [Dickson et al. 1992; Melnick et al. 1993].

The eggs laid by homozygous *Draf<sup>HMT</sup>* females [Fig. 1F] resemble strongly those laid by females homozygous for the EGF receptor mutation *top*<sup>1</sup> [Schupbach 1987; Clifford and Schupbach 1989], as shown in Figure 1C. The eggshells are ventralized, with fused and shortened dorsal appendages, and the eggs are longer than wild type. The chorions of ventralized eggs show an increased number of follicle cell imprints, with more follicle cells giving rise to the main body of the chorion at the expense of the dorsal appendages [Schupbach 1987]. The number of follicle cell imprints on eggs from *Draf<sup>HMT</sup>* mothers is ~15% higher than wild type (Table 1). Dorsoventral polarity can, therefore, be disrupted by mutations that affect either the activity or the levels of the EGF RTK or Raf, implying that these two kinases participate in a common pathway.

**Expression of constitutively activated Raf during oogenesis dorsalizes the eggshell**

Constitutive activation of the dorsoventral signaling pathway during oogenesis should lead to a dorsalized egg phenotype, the converse phenotype to that observed when signaling is reduced by mutations in the EGF RTK,
or in Draf. If Draf activity is sufficient for signaling, then ectopic expression of an activated form of the kinase should generate a dorsalized phenotype. Raf can be activated by deleting the amino terminus of the kinase, which is thought to encode a negative regulatory region (Stanton et al. 1989; Wasylyk et al. 1989; Bruder et al. 1992; Heidecker et al. 1992). Therefore, we expressed a constitutively activated form of either Drosophila or human Raf kinase in adult females and looked for an effect on oogenesis and embryonic development.

To create a heat-inducible, gain-of-function Raf gene (Raf°), we subcloned the coding sequence for amino-termi­nally deleted Drosophila (A. Brand, X. Lu, and N. Perrimon, in prep.) or human Raf (Heidecker et al. 1992) downstream of the hsp70 promoter. Raf° was expressed by heat shocking females carrying the hsp70--Raf° gene (see Materials and methods). After 3 days, in which heat shocks are administered once daily, the females lay eggs that are dorsalized and look similar to those laid by homozygous fs(1)K10 females (Fig. 1, cf. B with D). The dorsal appendages fuse in a ring at the anterior end, and the eggs are smaller and more spherical than wild type. A number of eggs exhibit a seemingly more severe phenotype where the dorsal appendages are reduced in size, or completely absent, and the chorion is deposited in a wide opaque band toward the anterior of the egg (Fig. 1E). The number of more severely affected eggs increases with the frequency and duration of the heat shock.

Higher levels of Raf, or increased kinase activity, may be responsible for the stronger chorion phenotype. This is supported by the result that expression of amino-termi­nally truncated human Raf, which appears to be more active than truncated Draf (A. Brand, X. Lu, and N. Perrimon, in prep.), gives rise to more eggs displaying the stronger phenotype. Expression of Draf° causes 63% of the eggs to become fs(1)K10-like, whereas 29% look more severe. By comparison, expression of human Raf° results in 55% resembling fs(1)K10-like eggs and 43% showing the more severe phenotype (Table 2). Females homozygous for the hsp70-human Raf° gene lay very few eggs after the third day of heat shock, of which two-thirds are severely dorsalized (Table 2). Ectopic activation of Raf may also affect an earlier stage of oogenesis, during which Raf plays an as yet unidentified role.

In contrast to ventralized chorions, dorsalized egg-shells bear fewer follicle cell imprints. Counting the fol­licle cell imprints on eggs laid by hsp70--Draf° females indicates 10% fewer imprints on the dorsalized chorions, and 25% fewer on the more severely affected eggs (see Table 1). Because the ectopic activation of either Draf or human Raf dorsalizes the egg, activating Raf appears to be sufficient to initiate dorsoventral signaling.

Table 2. Chorion phenotypes

| Genotype             | Total | Wild type (%) | Dorsalized (%) | More severe (%) |
|----------------------|-------|---------------|----------------|----------------|
| yy                   | 100   | 100 (100)     | 0              | 0              |
| hsp70--Draf°/TM3a    | 99    | 8 (8)         | 62 (63)        | 29 (29)        |
| hsp70-human Raf°/TM3a| 65    | 1 (2)         | 36 (55)        | 28 (43)        |
| hsp70-human Raf°     | 9     | 1 (11)        | 2 (22)         | 6 (67)         |
| UAS--Draf°/pGawB55Bb | 213   | 94 (44)       | 119 (56)       | 0              |
| UAS--human Raf°/pGawB55Bb | 144 | 5 (3)         | 139 (97)       | 0              |

*Eggs were collected from females that had been heat-shocked once daily for 3 days. Without heat shock, the females lay eggs with wild-type chorions.

In the absence of GAL4, UAS--Draf° and UAS--human Raf° females lay eggs with wild-type chorions.

Targeted expression demonstrates that Raf acts in the follicle cells to determine dorsoventral polarity

Expression of Raf° from a heat shock promoter cannot resolve whether Raf acts in the oocyte or in the follicle cells, as the kinase may be expressed both in the germ line and in the soma. To ask in which tissue Raf° has its effect, we targeted expression of activated Raf to the follicle cells using the GAL4 system (Brand and Perrimon 1993), which permits ectopic expression to be directed to particular cells or tissues. An enhancerless GAL4 gene is integrated randomly in the genome, generating independent insertion lines in which GAL4 expression is directed by different genomic enhancers. It is then possible to introduce a gene containing GAL4-binding sites within its promoter, to activate the gene in those cells where GAL4 is expressed, and to observe the effect of this directed misexpression on development.

To create a GAL4-responsive Raf° gene (UAS--Raf°), the human or Drosophila Raf° coding sequences were subcloned behind a tandem array of five optimized GAL4-binding sites (or UAS, for upstream activation sequence, see Materials and methods; Brand and Perrimon 1993). The UAS--Raf° gene is silent in the absence of GAL4. When GAL4 is introduced in a cross, GAL4 binds to the UAS and activates transcription.

We generated a library of 250 independent GAL4 insertion lines (Brand and Perrimon 1993) and characterized the GAL4 expression patterns by crossing them to a line carrying a UAS--lacZ gene and staining their progeny for β-galactosidase activity (Brand and Perrimon 1993; D. McKearin, pers. comm.) We also crossed the lines to a line carrying UAS--Raf°, and screened for Raf-dependent phenotypes.

One GAL4 insertion line, 55B, activates UAS--lacZ in a stripe of follicle cells around the anterior of the oocyte in a stage 9 egg chamber (Fig. 2A). Embryonic GAL4 ex-
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Figure 2. Targeted expression demonstrates that Raf acts in the follicle cells to determine the dorsoventral polarity of the egg. The dorsoventral polarity of the egg is established by signaling between the oocyte and the somatic follicle cells. As such, the polarity of the eggshell can be disrupted by mutations that affect either the germ line (such as $fs(1)K10$) or the follicle cells (such as $DER$). Using the GAL4 system to target gene expression (Brand and Perrimon 1993), we show that expression of activated Raf within the follicle cells leads to a dorsalized phenotype. The enhancer trap/GAL4 insertion 55B expresses GAL4 in the follicle cells of a stage 9 egg chamber, as shown in A where GAL4 insertion 55B drives expression of a UAS-lacZ gene. X-gal staining shows cytoplasmic $\beta$-galactosidase expression in a band of follicle cells surrounding the anterior end of the oocyte (marked by arrowheads). When line GAL4 insertion 55B is used to activate UAS-Draf or UAS-human Raf in the anterior follicle cells, the resultant eggs are dorsalized (B and C, respectively).

Figure 3. Activated Raf induces ventral follicle cells to adopt a dorsal cell fate. The dorsal anterior follicle cells can be marked by $\beta$-galactosidase expression from the enhancer trap/lacZ insertion AN296 (T. Schupbach, unpubl.), which carries an enhancer trap lacZ gene. The lacZ gene has integrated near a genomic enhancer that directs expression specifically in the dorsal anterior follicle cells [Figs. 3A, 4A, and 5A].

We expressed activated Raf from the hsp70-Draf gene in the AN296 background and assayed $\beta$-expression in ovaries. After three or four heat shocks, all the follicle cells over the oocyte in a stage 10 egg chamber express the previously restricted dorsal marker [Fig. 3B]. At stage 12 or 13, in a wild-type egg chamber, $\beta$-galactosidase expression is primarily restricted to the follicle cells that

Pression is restricted to the salivary glands. When line 55B is crossed to flies carrying UAS-Draf or UAS-human Raf, to express Raf specifically within the anterior follicle cells, the female progeny lay dorsalized eggs as judged by their chorion phenotypes [Fig. 2B,C; Table 2]. Activating Raf in the follicle cells is, therefore, sufficient to dorsalize the egg, suggesting that Raf is required in the follicle cells, rather than in the oocyte, to specify dorsoventral polarity.

Activated Raf induces ventral follicle cells to assume a dorsal cell fate

Expressing Raf in follicle cells might direct a cell fate switch, such that the ventral and lateral follicle cells assume a dorsal fate. To assay whether activated Raf can induce a cell fate change, we monitored the expression of a marker that is normally restricted to the dorsal anterior follicle cells. We used the transgenic line AN296 (T. Schupbach, unpubl.), which carries an enhancer trap lacZ gene. The lacZ gene has integrated near a genomic enhancer that directs expression specifically in the dorsal anterior follicle cells [Figs. 3A, 4A, and 5A].

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Figure 3. Activated Raf induces ventral follicle cells to adopt a dorsal cell fate. The dorsal anterior follicle cells can be marked by $\beta$-galactosidase expression from the enhancer trap/lacZ insertion AN296 (T. Schupbach, pers. comm.), as demonstrated by X-gal staining of an ovariole from a female carrying two copies of AN296. (A) In a stage 10 egg chamber only the follicle cells in the vicinity of the oocyte nucleus, which has migrated to a dorsoanterior position, stain. (B) In females that express Draf from a heat shock promoter, $\beta$-galactosidase expression from AN296 can be detected in all of the follicle cells overlying the oocyte.
Raf specifies dorsoventral polarity in Drosophila

Figure 4. Constitutive activation of Raf alters the morphology of the egg chamber. Stage 12 and 13 egg chambers from females expressing lacZ insertion AN296. [A] In wild-type animals, transcription of AN296 is primarily limited to the follicle cells that will give rise to the dorsal appendages. [B] In females expressing activated Draf° from a heat shock promoter, all of the follicle cells express AN296. [C] In some cases, a ring of densely packed follicle cells can be seen around the anterior end of the oocyte. These egg chambers may give rise to the most severely dorsalized eggs (see Fig. 1E).

Figure 5. Targeted expression of Raf°° dorsalizes the anterior follicle cells. Stage 10 egg chambers stained with X-gal to show: (A) β-Galactosidase expression from the enhancer trap/lacZ insertion, AN296, in a wild-type ovary and (B) targeted expression of activated human Raf, using the GAL4 insertion 55B, results in a high level of AN296 expression in a band of follicle cells surrounding the anterior end of the oocyte (cf. with Fig. 2A, showing the expression pattern of UAS–lacZ driven by GAL4 insertion 55B).

Expression of activated Raf dorsalizes the embryo as well as the chorion

We have shown that the ectopic activation of Raf is sufficient to dorsalize the chorion. Raf might act only to specify the fate of the dorsal follicle cells or, alternatively, localized kinase activation might also contribute to the dorsoventral polarity of the developing embryo. Therefore, we asked whether ectopic activation of Raf dorsalizes the developing embryo.

We prepared cuticles from embryos derived from females expressing Raf°° during oogenesis. Most of the eggs laid by females expressing activated Raf cannot be fertilized because of their dorsalized chorions (Tables 2 and 3). Of the fertilized eggs from females expressing UAS–human Raf°°, 25% give rise to dorsalized embryos (Fig. 6B). In the most severe cases, the embryos are twisted and lack nearly all ventral structures, such as the thoracic and abdominal denticle belts seen in a wild-type larva (Fig. 6A). More dorsally derived structures, such as the filzkörper, are still present (Fig. 6B). Embryos from hsp70–Draf°° females are comparably dorsalized. Fe-
Table 3. Cuticle phenotypes

| Genotype               | Σ  | Fertilized (%) | Wild type (%) | Dorsalized (%) | Weak* (%) |
|-----------------------|----|----------------|---------------|----------------|-----------|
| UAS-Draf<sup>red</sup>/pGawB<sup>55B</sup> | 150 | 126 (84)       | 101/126 (80)  | 4/126 (3)     | 21/126 (17) |
| UAS-human Raf<sup>red</sup>/pGawB<sup>55B</sup> | 117 | 28 (24)        | 11/28 (39)    | 7/28 (25)     | 10/28 (36)  |

*Weakly dorsalized and/or head defects.

males expressing UAS–Draf<sup>red</sup> produce far fewer dorsalized embryos (3%), suggesting that higher levels or activity of Raf are required to dorsalize the embryo as opposed to the chorion.

**Raf acts downstream of the EGF receptor**

We have shown that Raf acts in the follicle cells to relay a signal that establishes the dorsoventral polarity of the eggshell and the embryo. The phenotypes seen when Raf is ectopically activated, or when Raf activity is reduced, suggest that Raf might act downstream of the *Drosophila* EGF receptor. We examined the relationship between Raf and DER by expressing Raf<sup>red</sup> in a DER mutant background. If Raf acts downstream of DER, then activating Raf will overcome the need for a functional EGF RTK. The resultant eggs should therefore be dorsalized. If Raf acts upstream of DER, the absence of DER will suppress the dorsalized phenotype, and the eggs will be ventralized. If the two genes function in separate pathways, an intermediate phenotype might be obtained.

Using the GAL4 system, flies that are homozygous for the DER mutation, top<sup>1</sup>, can be made to express activated human or fly Raf in their follicle cells. Whereas top<sup>1</sup> females normally lay ventralized eggs (Fig. 7A), in the presence of constitutively active Draf (Fig. 7B) or human Raf (Fig. 7C) their egg shells are dorsalized. Similarly, expression of UAS–human Raf<sup>red</sup> in top<sup>1</sup>/top<sup>C1</sup>

![Figure 6](image1)

**Figure 6.** Activated Raf dorsalizes the embryo as well as the eggshell. Mutations that affect the dorsoventral pattern of the chorion can also alter the polarity of the embryo developing within the eggshell. When activated Raf is ectopically expressed, most eggs are dorsalized and as a result cannot be fertilized (76%, UAS–human Raf<sup>red</sup>). In those eggs that are fertilized, however, 61% of the embryos are dorsalized (A). A cuticle prepared from a wild-type larva [the vitelline membrane has been removed], showing the denticle bands on the ventral surface of the animal; (B) a cuticle prepared from an embryo laid by a female expressing UAS–human Raf<sup>red</sup> as directed by GAL4 insertion 55B. The embryo is twisted and lacks the ventral denticle bands, whereas the dorsally derived filzkörper is still present (arrowhead). In many embryos the dorsalization is more pronounced at the anterior end, as has been reported for *fs(1)R10* [Wieschaus 1979].

![Figure 7](image2)

**Figure 7.** Raf acts downstream of the EGF receptor. Females that are homozygous for mutations in the *Drosophila* EGF receptor (such as top<sup>QVY1/top<sup>QVY1</sup>) lay eggs that are ventralized, as shown in A. When Raf<sup>red</sup> is expressed in these females they lay dorsalized eggs, demonstrating that Raf<sup>red</sup> is epistatic to DER. Expression of either Draf<sup>red</sup> (B) or human Raf<sup>red</sup> (C, D) in the anterior follicle cells of females mutant for DER (B, C: top<sup>QVY1/top<sup>QVY1</sup>; D: top<sup>QVY1/top<sup>C1</sup>) dorsalizes the egg chamber.
Raf specifies dorsoventral polarity in Drosophila females results in dorsalization (Fig. 7D). Because expression of UAS–Draf° or UAS–human Raf° in the anterior dorsal and ventral follicle cells is sufficient to dorsalize eggs from DER mutant mothers, our results strongly suggest that Raf acts downstream of DER.

**Discussion**

Raf is a cytoplasmic serine/threonine that can transduce signals from several different receptor tyrosine kinases. In mammalian cells, Raf acts downstream of the platelet-derived growth factor and EGF RTKs (Morrison et al. 1988). During vulval induction in Caenorhabditis elegans a Raf homolog encoded by the lin-45 gene is required for signaling by the EGF RTK and Ras, encoded by let-23 and let-60, respectively (Aroian et al. 1990; Beitel et al. 1990; Han and Sternberg 1990; Han et al. 1993). In Drosophila, Raf has been shown to relay signals from the torso RTK to determine the fates of the embryonic termini (Ambrosio et al. 1989b), and from the sevenless RTK to specify the fate of the R7 photoreceptor cell (Dickson et al. 1992). Here we demonstrate a novel role for Raf in Drosophila development, in specifying the dorsoventral polarity of the egg chamber. We show that Raf functions in the somatic follicle cells to transduce a signal from the Drosophila EGF receptor (Fig. 8).

When Raf activity or expression levels are lowered by means of the hypomorphic mutation DraH, females lay ventralized eggs. The eggs are longer than wild type, have fused and shortened dorsal appendages, and exhibit more follicle cell imprints. The phenotype is presumably attributable to reduced signaling through the EGF receptor pathway. When constitutively active Raf kinase is expressed during oogenesis, the dorsoventral signal transduction pathway is ectopically activated, resulting in dorsalized eggs and embryos. Expression of activated Raf is sufficient to dorsalize both the chorion and the embryo, whereas the DraH mutation only ventralizes the chorion. The chorion may be a more sensitive indi-

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**Figure 8.** Raf transmits a signal from the EGF receptor to specify the dorsoventral polarity of the egg chamber. A diagrammatic representation of a stage 10 egg chamber. The region within the red box has been enlarged to show a segment of the dorsal side of the egg chamber. By this stage of development, the oocyte nucleus has migrated to the prospective dorsal side of the oocyte. The oocyte sends a signal to the dorsal follicle cells. This signal may be in the form of a ligand for the EGF receptor. Potential candidates for a ligand for DER include gurken and spitz. Activation of the EGF receptor initiates a signal transduction cascade that (1) determines the fate of the dorsal follicle cells and (2) establishes the polarity of the developing embryo. DraH seems to act downstream of the EGF receptor in both of these processes. In addition to Raf, several other signaling molecules have been shown to act downstream of the torso and sevenless RTKs. These include csw, Ras, and the MAP kinase activator (or MEK) encoded by Dsor1. Recent evidence suggests that these molecules also participate in dorsoventral signaling (L. Perkins, X. Lu, J. Duffy, and N. Perrimon, unpubl.).
cator of Raf activity than the embryo, such that ventralized embryos would only be seen after Raf levels were further reduced.

Raf can be activated by amino-terminal deletions that are thought to remove a negative regulatory domain. This model has been supported recently by work showing that the amino terminus of Raf interacts directly with Ras (Vojtek et al. 1993; Warne et al. 1993; Zhang et al. 1993). Ras may activate Raf by binding to the amino-terminal negative regulatory domain and releasing the carboxy-terminal kinase domain from negative regulation. A single amino acid substitution within the amino terminus of Draf, isolated as the hypomorphic mutation C110, may thus interfere with the Ras–Raf interaction and result in a less active kinase (Melnick et al. 1993; Vojtek et al. 1993). Expression of an amino-terminal region of Raf leads to a dominant negative phenotype in mammalian tissue culture cells (Bruder et al. 1992), suggesting that the amino terminus can compete with wild-type Raf for interaction with Ras.

When constitutively activated Raf is expressed throughout the egg chamber, the eggshell and the developing embryo are dorsalized. Therefore, although Raf is normally transcribed throughout the egg chamber (Ambrosio et al. 1989b), Raf activation appears to be restricted. To show that Raf acts in the somatic follicle cells rather than in the germ line, we targeted expression of Raf<sup>act</sup> to follicle cells using the GAL4 system (Brand and Perrimon 1993). The GAL4 system allows ectopic expression to be directed to particular cells or tissues. Expressing Raf<sup>act</sup> within the follicle cells alone dorsalizes the egg chamber. These results suggest that Raf is normally activated only within the dorsal follicle cells, and that activation within the ventral cells is sufficient to dorsalize the egg. We used GAL4 insertion line 55B to target expression of Raf<sup>act</sup> primarily to an anterior band of follicle cells. The levels of activation directed by GAL4 in this line appears to be somewhat variable from cell to cell, as judged by GAL4-mediated β-galactosidase expression. In spite of this, consistently we observe dorsalized eggs from expression of Raf<sup>act</sup> (Table 2), suggesting that the follicle cells are sensitive to small changes in the levels of Raf activity.

The dorsal signal emanating from the oocyte may form a gradient, and thus activate the EGF RTK pathway to different degrees depending on the concentration of ligand. This would be similar in principle to the the terminal class signal transduction pathway, where a ligand concentrated at the termini appears to result in a gradient of activation through the torso RTK (Casanova and Struhl 1989; Perkins et al. 1992; Melnick et al. 1993). High levels of torso activity lead to transcription of the genes tailless and huckebein, whereas lower levels of activity only support tailless transcription.

We were able to follow Raf<sup>act</sup>-induced cell fate changes by monitoring the expression of a dorsal follicle cell marker, the enhancer trap/lacZ insertion AN296. AN296 normally is expressed in a gradient radiating from the dorsal follicle cells. AN296 is expressed in all follicle cells in response to expression of hsp70–Raf<sup>act</sup>, and is transcribed strongly in a band of anterior follicle cells in response to targeted expression of UAS–Raf<sup>act</sup>.

To test whether Raf acts downstream of the EGF receptor, we expressed activated Raf in the dorsal and ventral follicle cells of DER mutant females. In the absence of activated Raf, DER mutants lay ventralized eggs. When activated Raf is expressed in their follicle cells, however, they lay dorsalized eggs. Because activating Raf overcomes the requirement for DER in specifying the dorsal follicle cell fate, Raf appears to act downstream of the EGF receptor. We have not ruled out the possibility that artificially activated Raf acts as a promiscuous kinase, and in this way bypasses the requirement for EGF receptor activation. However, the fact that loss-of-function and gain-of-function Raf mutants have opposite effects on dorsoventral signaling strongly suggests that Raf normally functions in the DER signal transduction pathway.

If the interaction domain within the amino terminus of Raf appears to be conserved between mammals, Drosophila and C. elegans (Han et al. 1993; Melnick et al. 1993; Vojtek et al. 1993) suggesting that Ras and Raf from different species might associate in a similar fashion. We show that expression of human Raf<sup>act</sup> can specify the dorsal follicle cell fate. Given the extent of conservation between many of the receptor tyrosine kinase signal transduction pathways, it seems likely that other members of the torso and sevenless pathways will be implicated in dorsoventral signaling. These include Ras1, Gap1 (a Ras GTPase-activating protein) and Sos (a guanine nucleotide exchange factor) acting upstream of Raf, csw (a tyrosine phosphatase) acting upstream of Raf or in a parallel pathway, and MEK and MAP kinase (Biggs and Zipursky 1992, Tsuda et al. 1993) acting downstream of Raf (Fig. 8). This is supported by recent results that suggest that Ras1, Gap1, csw, and Dso1 (the Drosophila MEK homolog) act in the dorsoventral pathway (Chou et al. 1993; X. Lu, L. Perkins, J. Duffy, and N. Perrimon, unpubl.).

Of the maternal effect genes that affect the dorsoventral polarity of both the egg and the developing embryo, only DER, thromboid, and Draf are required in the follicle cells. The factors downstream of DER, such as windbeutel, nudel, and pipe, affect only the embryonic dorsoventral axis, and are thought to generate the ligand for the receptor Toll, which specifies the ventral side of the embryo (Anderson et al. 1985; Hashimoto et al. 1988, 1991; Stein et al. 1991; Stein and Nüsslein-Volhard 1992). DER is proposed to have at least two roles during oogenesis: (1) to determine the dorsal follicle cell fate, and (2) to block the production of the ligand dorsally. Screening for enhancers and suppressors of the dorsalized phenotype produced by Raf<sup>act</sup> will be one means of identifying novel factors downstream of DER, those that affect the egg and the embryo, and those that affect only the embryo. It may then be possible to distinguish factors common to each of these processes, as well as to identify possible branch points in the two pathways. One target of DER and Raf appears to be the lacZ fusion gene in line AN296, which is transcribed in response to Raf
activation. The gene identified by this enhancer trap insertion might be involved in the morphological changes induced in the dorsal follicle cells, or could encode a negative regulator of windbeutel, nudel, and pipe.

**Materials and methods**

**Drosophila strains**

Flies were raised on standard *Drosophila* media at 25°C, unless otherwise noted. Descriptions of balancers and mutations that are not described in the text can be found in Lindsley and Zimm (1992).

The *Draf* allele used in this study is *Draf^PM7* (Kramers et al. 1983; Melnick et al. 1993). *Draf^PM7* fails to complement the *Draf* deficiency *Df(1)64C18*, and can be rescued by a P-transposable element carrying the *Draf* gene contained within a 4.3-kb genomic BamHI fragment (Nishida et al. 1988). The *Draf^PM7* gene has been cloned and sequenced, but no amino acid change has been found within the coding region, suggesting that the mutation may reside within transcriptional regulatory sequences and result in reduced levels of *Draf* expression (Melnick et al. 1993).

The EGFR mutations, *top^1* and *top^2/^fl+*, were kindly provided by T. Schupbach, and are described in detail by Schupbach (1987) and Clifford and Schupbach (1989). *fs(1)K10^+* was kindly provided by E. Wieschaus and has been previously described (Wieschaus et al. 1978).

The enhancer trap *lacZ* line, AN296 (T. Schupbach, unpubl.), carries the P-transposable element described by Bier et al. (1989) and was a kind gift from T. Schupbach.

**Gene fusions**

A cDNA encoding the human Raf kinase, deleted between amino acids 2 and 334, [Heidecker et al. 1992] was excised as a BamHI to *XbaI* fragment from plasmid pRafBB (a gift from N. Williams) and subcloned into the *BglII* and *XbaI* sites of P-element vector pCaSpeR-hs (described by Thummel and Pirrotta 1992) to make phsAcRaf.

The coding sequence for an amino-terminally truncated version of *Draf* was subcloned as an *EcoRI* to *XbaI* fragment from vector pUAS-*Draf* (A. Brand, X. Lu, and N. Perrimon, in prep.) into *EcoRI* and *XbaI* cut pCaSpeR-hs, to make phs*ΔDraf*. The *Draf*-coding sequence is deleted between amino acids 2 and 343, based on the amino acid sequence reported by Melnick et al. (1993).

To create a gene under the control of the yeast transcriptional activator GAL4, a cDNA encoding the truncated version of human Raf-1 (Δ2-334) was subcloned as an *EcoRI*-*XbaI* fragment into vector pUAST [Brand and Perrimon 1993] to give pUAS-*ΔRaf1*.

**P-element transformation**

Transgenic lines were generated by injection of CsCl banded DNA, at a concentration of 600 μg/ml, into embryos of strain *yw*, +/+, *Sc*, *P[ry +, Δ2-3]/TM6, Ubx* (Robertson et al. 1988) using standard procedures (Spradling 1986). Several independent transformants were obtained for each construct. In this study we use lines *hsac-Raf1^4-1/^TM3* or *hsac-Raf1^4-1/^hsac-Raf1^4-1+* to express *hsRaf1-human Raf^αβ*(αβ) and *hsDraf^αβ/^TM3* (A. Brand, X. Lu, and N. Perrimon, in prep.) to express *hsRaf1-human Raf^αβ* for GAL4-mediated expression of *UAS-human Rafαβ*.

The *GAL4* system

To express activated Raf in ovaries, we used a novel system for targeting gene expression [the *GAL4* system; Brand and Perrimon 1993]. An enhancerless gene encoding the yeast transcriptional activator GAL4 is inserted randomly into the *Drosophila* genome where, depending on the site of integration, expression is directed by any one of a diverse array of genomic enhancers. A second gene, containing GAL4-binding sites within its promoter, can then be introduced into this background where it will only be transcribed in those cells where GAL4 is expressed.

An enhancer detection screen carried out to recover lines that express GAL4 in a cell- or tissue-specific manner (Brand and Perrimon 1993). Each of the 220 independent GAL4 insertion lines we recovered was crossed to a line carrying the *UAS-lacZ* reporter gene (*UAS-lacZ^+1-2*; Brand and Perrimon 1993). Embryos from the cross were stained for β-galactosidase expression with anti-β-galactosidase antibodies. The lines were then crossed to line *UAS-ΔRaf1^αβ* to screen for Raf^αβ*-dependent adult phenotypes.

pGawB^58^

Females derived from a cross between GAL4 insertion line, pGawB^58^, and *UAS-ΔRaf1^αβ* survive to adulthood and lay dorylized eggs. pGawB^58^ carries an enhancer trap/GAL4 vector (pGawB; Brand and Perrimon 1993) inserted on the third chromosome.

To characterize the ovarian GAL4 expression pattern, GAL4-directed expression of β-galactosidase was assayed. Line pGawB^58^ was crossed to line *UAS-lacZ^+1-2* (Brand and Perrimon 1993), and the progeny of the cross were allowed to develop to adults. Females were fed for 3–5 days, and ovaries were then dissected and stained for β-galactosidase activity, as described below.

**Staining ovaries for expression of β-galactosidase**

Ovaries were dissected in PBS, 0.1% Triton X-100 and fixed in 1% glutaraldehyde for 15–20 min. They were then washed in PBS, 0.1% Triton X-100 and stained in a solution of 10 mM NaH₂PO₄, 150 mM NaCl, 3 mM MgCl₂, 0.5 mM K₃[Fe(CN)₆], 3 mM K₄[Fe(CN)₆] containing a 1:50 dilution of X-GAL (25 mg/ml in dimethyl formamide). Ovaries were stained at room temperature for periods ranging from 1 hr to overnight. After washing in PBS, 0.1% Triton X-100, ovaries were mounted in 70% glycerol.

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Mounting eggs and cuticles

Eggs were mounted in a 1:1 mixture of Hoyer’s mountant and lactic acid, and then cleared overnight on a hot plate at 65°C. To highlight the follicle cell imprints, chorions were photographed using a 10X phase contrast objective with the condenser set to the phase 3 position. Follicle cell imprints were counted on an enlarged photograph showing a lateral view of each chorion. The counts are, therefore, very approximate, and are not comparable to those given by Schupbach (1987).

Cuticles were prepared and mounted as described by Struhl (1989).

Epistasis experiments

Eggs laid by females homozygous for the EGF receptor mutation top^f1 were compared with eggs laid by top^f1 females expressing Raf^f0. The GAL4 system was used to express either UAS–Drafl^f0 [in females of the genotype top^f1/top^f1, pGawB^55B/UA5–ΔDraf^f179] or UAS–human raf^f0 [in females of the genotype UAS–ΔCraf^f102/+; top^f1/top^f1, pGawB^55B/+ and UAS–ΔCraf^f102/+; top^f1/top^f1; pGawB^55B/+; see Fig. 7, C and D].

Acknowledgments

We thank Joe Duffy for discussions throughout the course of this work, for communicating results before publication, and for comments on the manuscript. We are most grateful to Trudi Schupbach for providing numerous stocks, including her unpublished line AN296, and for helpful discussions. Thanks go to Daniel St. Johnston for his drawing of the stage 10 egg chamber shown in Figure 7, and to Beth Noll for developing and printing most of the black-and-white photographs. We thank Xiangyi Lu, Eric Wieschaus, Linda Ambrosio, Carl Thummel, and Vincent Pirrotta for providing Drosophila strains or DNAs, and Trudi Schupbach, Dennis McKearin, Xiangyi Lu, Tze-Bin Chou, and Liz Perkins for communicating results before publication. Many thanks to Daniel St. Johnston, Acaimo Gonzales-Reyes, Jonathon Pines, and Jim Haseloff for comments on the manuscript. A.B. was supported by postdoctoral fellowships from the Helen Hay Whitney Foundation and the National Institutes of Health, and by a Leukemia Society of America Special Fellowship. This work was supported by the Howard Hughes Medical Institute.

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Raf acts downstream of the EGF receptor to determine dorsoventral polarity during Drosophila oogenesis.

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Genes Dev. 1994, 8:
Access the most recent version at doi:10.1101/gad.8.5.629

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