The transcription factor CF2 is a mediator of EGF-R-activated dorsoventral patterning in Drosophila oogenesis

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Establishment of dorsoventral polarity during Drosophila oogenesis requires localized intercellular communication between the follicular cells and the oocyte. This is initiated by the transmission of a “dorsal signal” from the oocyte to the anterior dorsal follicle cells by the EGF receptor (EGF-R) pathway and is followed by transmission of a second signal from the ventral follicle cells back to the embryo. We show that the zinc finger transcription factor CF2 participates in these processes. CF2 is suppressed by EGF-R signaling in the anterior dorsal follicle cells. Altered expression patterns of CF2 result in specific dorsoventral patterning defects in egg chambers and in embryos, as demonstrated phenotypically and with molecular markers. CF2 appears to act as a repressor of dorsal follicle cell fates and specifically as a repressor of the rhomboid gene transcription.

[Key Words: Dorsoventral patterning, EGF receptor signaling, oogenesis, transcription repression]

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Dorsoventral (D/V) pattern in the Drosophila egg chamber, which also affects subsequent embryogenesis, is generated through the action of ≈20 maternal-effect genes (for reviews, see Schüpbach et al. 1991; Chasan and Anderson 1993). This pattern is initiated at stages 8–9 of oogenesis, when the oocyte nucleus migrates to an anterodorsal position. Soon afterward gerken (grk) RNA and the protein it encodes, a transforming growth factor α (TGFα)-like ligand, become similarly localized in the anterodorsal region of the oocyte neighboring the nucleus (Neuman-Silberberg and Schüpbach 1993; González-Reyes et al. 1995; Roth et al. 1995). Localization of grk RNA is the critical step and is ensured by at least three mechanisms: (1) mRNA synthesis in the dorsally positioned nucleus; (2) action of the fs(1)K10 (henceforth K10) gene product probably to reduce transcription (Prost et al. 1988; Forlani et al. 1993); and (3) prevention of mRNA diffusion to the ventral side through the actions of at least four genes: spirale (spir), cappuccino (capu), orb, and squid (squil) (Manseau and Schüpbach 1989; Kelly 1993; Neuman-Silberberg and Schüpbach 1993; Christenson and McKearin 1994; Roth and Schüpbach 1994).

The spatially localized Grk protein activates in the facing anterior dorsal follicle cells the membrane-bound, ubiquitous epidermal growth factor receptor (EGF-R) that is encoded by the torpedo (top) locus (Price et al. 1989). Grk binding initiates the EGF-R signal transduction pathway, which involves homologs of Ras, Raf, and MEK (Brand and Perrimon 1994; Hsu and Perrimon 1994; for review, see Perrimon 1994), and ultimately establishes the dorsal fate of the follicular cells. In mutants such as K10 and spir, the grk RNA and presumably the Grk protein spread to the ventral side of the oocyte, resulting in dorsalized egg chambers (Neuman-Silberberg and Schüpbach 1993; Roth and Schüpbach 1994). Dorsal fate is manifested by specialized chorion structures elaborated by the follicular cells, such as the respiratory dorsal appendages (Margaritis et al. 1980). Another transmembrane protein, encoded by rhomboid (tho), is expressed in the follicle cells where top is activated and probably serves to amplify the dorsal signal (Ruohola-Baker et al. 1993).

In contrast, when gene functions such as grk, top, Draf, and tho are absent, dorsal cell fates are insufficiently defined and the egg chamber becomes ventralized (Schüpbach 1987; Ruohola-Baker et al. 1993; Brand and Perrimon 1994). These mutations, as well as K10 and spir, also result in corresponding embryonic phenotypes, indicating that the D/V patterning processes in the egg chamber and embryo are coupled, through the EGF-R signaling pathway (Schüpbach and Wieschaus 1989, 1991).

Because the Grk ligand is localized dorsally, the ventral follicle cells do not normally experience EGF-R sig-
nalizing. Instead, they undergo a ventral-specific signaling process that activates ultimately an embryonic membrane receptor, the Toll gene product [Hashimoto et al. 1991; Schneider et al. 1991]. Toll activation results in translocation of the transcription factor Dorsal into the nuclei of the syncytial blastoderm, in a gradient that specified localized expression of different sets of zygotic genes [e.g., twist (twi) ventrally and decapentaplegic (dpp) dorsally], ultimately leading to D/V differentiation of embryonic structures [Jiang et al. 1991].

Although many components of these pathways have been described, an important gap is that no transcriptional regulators have been identified in follicular cells, which are either controlled by the dorsal signal or themselves control ventral signal production. It is likely that transcription factors that are critical for the maternal D/V patterning processes are also essential for other events in embryonic and postembryonic development [Ruohola-Baker et al. 1994]. Mutations in these factors might result in lethality, precluding detection of their effects on oogenesis, which occurs at a late stage of the life cycle. We examined a collection of transcription factors that are expressed in the ovaries [Shea et al. 1990], and noted that a nuclear zinc finger protein CF2 [Hsu et al. 1992] shows spatially restricted expression in the follicular epithelium, suggesting a role in spatial patterning. Here we demonstrate that indeed CF2 is involved in this process in a manner that may help explain the development of both follicular and embryonic D/V patterning.

Results

**Expression of the transcription factor CF2 is regulated spatially by the EGF-R pathway**

CF2 is a Cys$_2$-His$_2$, zinc finger protein defined initially by virtue of its sequence-specific binding to the promoter of the chorion [eggshell] protein gene $s15$ [Shea et al. 1990]. Although its functional role in chorion gene expression has not been demonstrated in vivo, immunocytochemistry identified CF2 in the nuclei of follicle cells that surround the oocyte and produce the chorion. Interestingly, CF2 protein is detected initially at early stage 8 of oogenesis in all the follicle cells that surround the oocyte, but later becomes spatially restricted; at late stage 9 and early stage 10 it disappears from the anterodorsal follicle cells, creating a prominent negative “patch” at least through stage 12 [arrow in Fig. 1, left column].

The patch corresponds to the region where the oocyte and the somatic follicle cells communicate through the EGF-R signaling pathway, resulting in the determination of dorsal follicle cell fates. We showed that mutations involving this pathway affect the localized depletion of CF2; the CF2-negative patch expands into the ventral region in the dorsalized mutants sprt and especially $R10$. in pace with the expansion of dorsal cell fate [Fig. 1, middle column]. Conversely, the patch shrinks in ventralized mutants and is completely eliminated in strong mutants of grk and top [Fig. 1, right column].

**CF2 protein functions as a repressor of dorsally expressed genes**

The observations indicated that CF2 is depleted as a result of EGF-R signaling. Manipulation of CF2 levels indicated further that CF2 can quickly repress [directly or indirectly] gene activities that are associated with the dorsal cell fate; rho, which is first induced at stage 10 and is required for the dorsal fates of the anterodorsal cells [Ruohola-Baker et al. 1993; Brand and Perrimon 1994], and the enhancer trap insert in line AN296 [Bier et al. 1989], which responds in a cell-autonomous fashion to alterations in the activity of the EGF-R pathway [J. Duffy and N. Perrimon, pers. comm.]. For these experiments we used CF2 sense and antisense constructs directed by an hsp70 promoter (henceforth CF2-s and CF2-as, respectively); we confirmed that CF2-s induction results in the presence of CF2 in all follicular cells, whereas CF2-as induction depletes CF2 in ventral as well as dorsal cells, especially in the anterior region [Fig. 1, lower right and middle]. As shown in Figure 2, at stage 10 and within 2 hr of the heat shock, overexpression of CF2 (CF2-s) represses both AN296 and rho expression in the anterior dorsal follicle cells, whereas depletion of CF2 (CF2-as) causes AN296 expression to spread ventrally and posteriorly, and rho expression to become generalized in the follicular epithelium. At stage 11, normal rho expression consists of two elements: a thin anterior layer of cells, between the nurse cells and the oocyte, and a V-shaped “double stripe”, corresponding to the dorsal appendages [Ruohola-Baker et al. 1993]. The former remains immediately susceptible to CF2 repression but, interestingly, the double stripe loses susceptibility to CF2 once the dorsal fate has been established at stage 10 [Fig. 2]. If CF2-s is induced 10 hr earlier [i.e., at stage 9–10a], the double stripe at stage 11 is suppressed [data not shown].

**Ectopic expression or depletion of CF2 alter the D/V polarity of the egg chamber**

We examined whether ectopic overexpression or depletion of CF2 during oogenesis might alter D/V polarity of the egg chamber. Such effects were seen, which for simplicity will be summarized as dorsalization or ventralization, respectively [see also Discussion].

In the top and grk ventralized mutants, egg chambers become elongated in shape, their dorsal appendages fuse and shrink, and the number of main body follicle cells increases [Fig. 3; Table 1]. When CF2 protein was overexpressed from two copies of the CF2-s transgene, a high proportion of the egg chambers showed at least moderate ventralization, in all three respects [CF2-s; Fig. 3; Table 1]. These phenotypes were attributable to CF2 overexpression, as they were almost never seen in the identically heat-shocked parental strain [yw; Figs. 3 and 4] or in the same CF2-s strain without heat shock [results not shown]. At least two transgenic lines were used in these and subsequent experiments, with consistent results.
Transcription factor in dorsoventral patterning

Figure 1. CF2 protein distribution in wild-type and dorsoventral polarity mutant egg chambers. All egg chambers were stained with CF2 monoclonal antibody and examined laterally by confocal microscopy. Anterior is to the left and dorsal side is up. Phenotypes are indicated at the top of the three columns and genotypes in individual panels. Note that CF2 is mostly nuclear. Arrows point to areas in the oocyte-associated follicle cell layer where CF2 expression is depleted. Panels in the top row show a surface view and an optical cross section of the same egg chamber. All other egg chambers except the wild type at stage 12 (bottom left) are shown in optical cross section. In the wild type, CF2 appears at early stage 8 in the oocyte-associated follicle cells (bottom left), but by early stage 10, CF2 it disappears from an anterodorsal patch of follicle cells (top left); this feature persists at least through stage 12 (bottom left). In dorsalized mutants, the CF2-depleted patch is more extensive; in KIO (top middle), it spreads ventrally to become a distinct band surrounding the anterior region of the oocyte, but in spii the boundary between CF2 expressing and nonexpressing cells is less well defined (middle). Similar spreading of the patch results by inducing high levels of CF2 antisense RNA from an hsp70 gene promoter (CF2-as, bottom middle). In the ventralized mutants gsk (top right) and top (middle right), the patch lacking CF2 expression is absent. Similar restoration of CF2 expression can be obtained by inducing two copies of the hsp70–CF2 transgene (bottom right); in this case CF2 protein is also detectable in the normally unstained border cells and the squamous follicle cells associated with the nurse cells (anterior).

Conversely, when four copies of the CF2-as construct were induced by heat shock, phenocopies were observed resembling moderately although not extremely dorsalized mutants [like spii but not KIO in Fig. 3]. The egg chambers assumed a more rounded shape with enlarged and fused appendages and reduced follicle cell counts (Fig. 3; Table 1).

To establish at what stage heat-induced changes in CF2 expression trigger phenocopies, we examined eggs laid at different times after a pulsed heat shock. Using an oogenesis time scale (Spradling 1993), we could infer from the time delay at what stages of oogenesis the analyzed eggs had been exposed to CF2-s or CF2-as action. The data are shown in Figure 4. Both dorsalized and ventralized phenocopies were observed resembling moderately although not extremely dorsalized mutants [like spii but not KIO in Fig. 3].

Figure 2. CF2 negatively regulates dorsal follicle cell-specific genes. Ovaries were dissected from well-conditioned females 1.5 hr after completion of pulse heat treatment and were immediately processed for X-gal staining or RNA hybridization. (Left column) Lateral views of X-gal stained stage 10 egg chambers from the enhancer trap line AN296 in the three genetic backgrounds indicated at left. Expression of the lacZ marker directed by AN296 is indicated by arrows. Note the complete elimination of AN296 expression in CF2-s and its ventral expansion in CF2-as. The other columns show dorsal or dorso-lateral views of egg chambers of the same three genotypes, in situ hybridized for rho RNA; two oogenic stages, 10 and 11, are shown (indicated at top). The area of wild-type stage 10 rho expression in the oocyte-associated follicle cell layer is marked by a bracket. At stage 10 (middle column), ectopic CF2 (CF2-s) represses rho expression, whereas antisense RNA (CF2-as) results in ectopic rho expression throughout the egg chamber. At stage 11 (right column), the double stripe element of rho expression on the dorsal surface is not affected by CF2-s or CF2-as, but expression in a thin anterior layer of follicle cells bordering the nurse cells and the oocyte (arrowheads) is repressed in CF2-s.
Mosaic analysis suggests that loss of CF2 can lead to overproduction of dorsal appendage material

We have used mitotic clone analysis to obtain independent genetic evidence suggesting that CF2 depletion may promote a specific dorsal cell fate, at least in anterior dorsal follicle cells that are exposed to EGFR signaling. Point mutants of CF2 are not currently available. However, Df(2L)27 is a small (~25 kb) deletion, null for CF2, which removes the entire CF2-coding region and DNA 5' to it [J. Tower, pers. comm., C. Bagni, S. Bolshakov, and C. Mollinari, unpubl.]. Although we cannot exclude the possibility that genes outside the limits of the deficiency might also be affected, no ovarian transcripts other than CF2 have been detected using as probe DNA that is encompassed by this deficiency (data not shown).

Rare defects that apparently represent clonal expansion of a specific type of dorsal fate were observed in eggs.
laid by deficiency heterozygotes that had been X-irradiated either as adults (8/5000; 0.16%) or as first instar larvae (60/6700; 0.9%). These defects were never observed in identically irradiated \textit{yw} controls. Although the chromosomes were not marked, homozygous clones resulting from X-ray-induced mitotic recombination can be expected at similar frequencies (Wieschaus and Gehring 1976; Stevens et al. 1990). Scanning electron micrographs of these defects are shown in Figure 5. In the normal chorion (Fig. 5A; Margaritis et al. 1980), the anterior dorsal domain bears two prominent and separate respiratory appendages, smooth near their base and fenestrated more distally to form a respiratory plastron (Spradling 1993). The rest of the chorion surface is marked with polygonal imprints of secretory follicular cells, which are seen in greater relief between the two appendages and in the extreme anterior operculum. Defects seen after X-irradiation of deficiency heterozygotes involve a substantial excess of smooth material resembling the base of the dorsal appendages. This material can replace the various kinds of cell imprints in the operculum, between the appendages or in large sectors of the main body, often it follows polygonal borders, suggestive of clonal origin (Fig. 5E). In other cases, the extra material forms multiple knob-like protrusions (Fig. 5C,D) or even short supernumerary appendages (Fig. 5B). Fenestrated plastron-like surfaces appear ectopically (Fig. 5F).

\textit{CF2} can also direct embryonic D/V patterning through a maternal effect

Having shown that changes in the level of \textit{CF2} during oogenesis can alter the D/V pattern of the egg chamber, we examined whether it may similarly affect the D/V polarity of the developing embryo. At 14–19 hr after \textit{CF2}-s induction, flies laid a significant number (46%) of eggs that developed into nonviable embryos showing a severely ventralized phenotype with expanded ventral denticles; the phenotype was similar to that of \textit{top} mutants (Fig. 6A, cf. \textit{top} and \textit{CF2}-s). In contrast, similar treatment of \textit{CF2-as} flies resulted in a significant number (29%) of dorsalized embryos that essentially lacked ventral denticles but showed a characteristic “twist” in the anterior cuticle; they resembled mutant embryos laid by \textit{KIO} females, except that in \textit{KIO} the posterior segment is often ventralized (Roth and Schüpbach 1994), whereas \textit{CF2-as} induces dorsalization throughout the ventral epidermis (Fig. 6A, cf. \textit{KIO} and \textit{CF2-as}). Because the eggs were laid 14–19 hr after fly heat treatment, the embryonic phenocopies resulted from maternal misex-

Figure 5. X-ray-induced mitotic clones of \textit{Df(2L)y\textsuperscript{2}} cause an expansion of dorsal follicle cell fate. The chorion surfaces of laid eggs were examined by scanning electron microscopy; the anterior end is at the top. First-instar larvae, heterozygous for the \textit{CF2} null \textit{Df(2L)y\textsuperscript{2}} deletion, were irradiated and the resulting adult females produced some defective eggs as shown. (A) A wild-type egg exhibits clear separation of the two dorsal respiratory appendages (solid arrowhead), each consisting of a smooth shaft (arrow) and a more distal fenestrated plastron (open arrowhead). Also note the follicle cell imprints in the chorion, which are most prominent between the appendages and in the anterior operculum cap, the pointed micropyle and the horse-shoe-shaped curved collar. (B) Two short supernumerary appendages (arrows), presumably arising from small clones of cells at the base of the normal appendages. The surrounding follicle cell imprints are normal. The right outgrowth has a plastron-like surface (inset). (C) An extensive area is covered by smooth dorsal material that resembles the shaft of the dorsal appendages, interspersed with plastron-like structures. Note the knob-like bumps and protrusions, and the posterior extension (arrowhead) of this putative clone. (D) An even larger clone affecting most of the left side of the egg has caused a partial loss of the collar (arrowhead) and deposition of thick dorsal material covering the operculum. Interestingly, a second collar-like structure is evident on the right. (E) The expansion of dorsal fate appears clonal. The thick smooth material follows exactly the polygonal cell outlines left by the follicle cells (arrowhead). In a closeup of the same egg (F), smooth dorsal material replaces the normally prominent cell imprints between the appendages (open arrowhead) and on the operculum (closed arrowhead). A few cell imprints showing plastron-like material are evident (arrow).
expression of CF2 at stages 9–10a of oogenesis. These phenocopies were not observed in the progeny of heat-shocked wild-type flies or of transformants that were not heat-shocked [data not shown].

At comparable frequencies, defects were seen in the expression of dpp and twi (dorsal and ventral markers, respectively; St. Johnston and Gelbart 1987; Thisse et al. 1988) in embryos laid by heat-shocked flies with altered levels of CF2 [Fig. 6B]. Follicular expression of the CF2-as construct led to molecular dorsalization of ≈26% of the embryos; the affected blastoderms showed generalized expression of dpp, superimposed on the normal dorsal expression, and essentially complete lack of expression of the ventral marker twi. In contrast, expression of CF2-s essentially abolished dpp expression and led to generalized twi expression, superimposed on the normal ventral domain of twi in ≈36% of the embryos.

Ectopic expression of CF2 counteracts the dorsalizing effects of K10 mutants

Dorsalizing mutations such as K10 spread the negative CF2 patch toward the ventral side of the follicular cell layer (see Fig. 1). An important test of the hypothesis...
that CF2 is functionally downstream of these genes is that the effects of CF2 misexpression should be epistatic, overexpression of CF2 in *K10* backgrounds should counteract the dorsalized chorion and embryonic phenotypes. We confirmed by immunostaining that, as in the wildtype, heat shock induces the CF2-s transgene to produce CF2 throughout the *K10* follicular epithelium [data not shown].

CF2 overexpression can reduce the dorsalizing effects of *K10*, rescuing ~10% of the egg chambers (10 of 103 examined), to the point that they showed two distinct rather than fused appendages [see Fig. 3, rescued *K10*]. Similarly, CF2 overexpression during oogenesis can reverse the maternal dorsalizing effect of the *K10* mutant on embryonic development [Fig. 6A, rescued *K10*]. About 12% of the embryos (12 of 109) derived from *K10; CF2-s* had the ventral denticles restored extensively, yielding a phenotype outside the range normally observed in embryos from *K10* mothers. Similar rescues were observed with the weaker dorsalizing mutant spir*RF48*.

**Ectopic expression of CF2 exacerbates heterozygous top mutant phenotypes**

Contrary to the partial rescue of dorsalized phenotypes, ectopic expression of CF2 is expected to exacerbate ventralizing mutants such as *top*. This is best illustrated in the heterozygous *top* background, which produces largely wild-type egg chambers with only 2% ventralized. As shown in Table 2, a mild heat treatment (3 days at 30°C of heterozygous *top* flies has no phenotypic effect, but in the presence of *CF2-s* [*top/SM6; CF2-s*] it generates 43.4% ventralized eggs, a percentage that is also significantly higher than *CF2-s* alone (31.6%). With more intense heat treatment [30°C for 3 days plus two incubations at 37°C for 15 min daily], the frequency of ventralized eggs in *top/SM6* alone remains very low (3%), but in *top/SM6; CF2-s* it reaches 100%, a severity the same as in homozygous *top* and again significantly higher than *CF2-s* alone (81.8%).

**Discussion**

**CF2 as a component of the D/V patterning pathway**

We have presented evidence that in the aggregate, strongly implicates CF2 in D/V patterning during oogenesis. Our interpretation is that CF2 is one of probably several transcription factors that act in concert, within specific domains of the follicular epithelium and in response to the EGF-R tyrosine kinase (RTK) signaling pathway, to pattern the epithelium and the eggshell that it produces, and indirectly to define the D/V axis of the embryo. The evidence is as follows:

1. CF2 protein expression is down-regulated in the critical anterodorsal domain (patch) of the epithelium, where the dorsal signal is received. Suppression of CF2 begins around the time that the signal emanates from the oocyte, at stage 9–10.
2. Suppression of CF2 is under the control of the genetic hierarchy that establishes the dorsal and ventral fol-

**Table 2:** Overexpression of CF2 exacerbates the heterozygous top mutant

| Genotypes | 25°C | 30°C | 30°C + 37°C |
|-----------|------|------|------------|
| *yw*      | 0    | 0.5  | 1          |
| *top/top* | 100  | 100  | 100        |
| *top/SM6* | 2    | 2    | 3          |
| *CF2-s*   | 1    | 31.6 | 81.8       |
| *top/SM6;CF2-s* | 3.5 | 43.4 | 100        |

All fly strains carry *yw* markers.

*Young female flies conditioned previously for 3–4 days at 25°C were dissected for ovaries after a 3-day incubation period at the designated temperatures. 30°C + 37°C denotes additional heat shock of 15 min twice daily at 37°C in a 3-day 30°C incubation period.

1. In each condition, ovaries were dissected at the end of incubation, and 200–250 stage 14 eggs were scored.

licular cell fates. In particular, it is responsive to the maternal-effect genes *K10, spir, grk,* and *top.*

3. CF2 itself can alter D/V patterning. In particular, ectopic depletion of CF2 spatially expands the expression of the dorsal gene markers AN296 and rho. Conversely, ubiquitous CF2 overexpression [presumably the persistence of CF2 in the anterodorsal follicular cell patch] has the opposite effect, blocking the expression of these dorsal genes.

4. Significantly, CF2 depletion and CF2 overexpression produce opposite effects, which resemble partial dorsalization and ventralization of the anterior chorion, respectively. Moreover, changes in CF2 during oogenesis have long-term effects; depletion or overexpression result in dorsalization or ventralization of the embryo, respectively, subsequent to fertilization, according to both cuticular and molecular markers.

5. Specific and spectacular effects on chorion structure are observed by X-irradiation of flies heterozygous for a small deficiency, null for CF2. The defects are localized, with irregular borders that follow cell imprint borders, and apparently correspond to overexpression of dorsal appendage material, sometimes in large sectors that can encompass the operculum and major parts of the main body, often they result in formation of extra appendages or appendage-like knobs.

6. CF2 overexpressed during oogenesis dramatically counteracts the dorsalizing effects of the *K10* mutant, both on the follicular chorion product and on the postfertilization embryo. Conversely, CF2 overexpression exacerbates a top heterozygote, which by itself produces wild-type chorions. Thus, CF2 is downstream of the well-known determinants of D/V polarity, *K10,* and *top.*

We have not demonstrated rigorously that the observed effects are somatic [follicular] cell specific, rather than mediated by the germ-line cells [nurse cells and oocyte]. However, this conclusion is supported by the evident EGF-R-controlled patterns of CF2 expression in
the follicular cell, by our inability to detect CF2 in the oocyte even upon CF2-s induction, and by the lack of immunostainable CF2 in embryos earlier than stage 13 (9 hr; C. Bagni, J.A. Gogos, S. Bray, and F.C. Kafatos, in prep.). The observed D/V defects resulting from changes in CF2 levels were described as pattern dorsalization and ventralization; they resemble the defects associated with weak to moderate mutations high on the D/V hierarchy but are not as extreme as those associated with the strongest alleles of K10 and grk. This might be expected for heat shock-driven constructs, especially for antisense constructs that are often only partially effective. On the other hand, some differences appear to be more than quantitative, suggesting that CF2 is only one of several transcription factors mediating the effects of the EGF-R signaling pathway, and is somewhat specialized in its effects.

**The mechanism of CF2 action in D/V patterning**

On the basis of our results and previous information, we present in Figure 7 a plausible model for CF2 action.

From the mutant phenotypes and the fact that immunostainable CF2 begins to disappear from the anterodorsal patch at late stage 9, before rho RNA appears in the same cells at early stage 10, we infer that activation of EGF-R by Gurken binding clears away CF2 through a mechanism that initially does not require rho. This is consistent with the observation that initially the overexpressed CF2 acts upstream of rho to repress the early rho expression pattern. In the normal course of events, suppression of CF2 permits expression of rho, AN296, and other aspects of the dorsal cell fate, including formation of normal dorsal appendages. However, we do not know yet whether the apparent repressor function of CF2 is direct or indirect.

We have not investigated the steps between activation of EGF-R and suppression of CF2. However, from previous studies on ovarian patterning we know that activated EGF-R and Rho act in concert to set in train a signal transduction cascade that includes the Ras, Raf, and MAPK kinase (MEK) protein kinases (Ruohola-Baker et al. 1993; Brand and Perrimon 1994; Hsu and Perrimon 1994; Lu et al. 1994). This cascade may also include MEK-activated mitogen-activated protein kinase (MAPK), which is known to translocate into the nucleus and phosphorylate diverse transcription factors in vertebrates and certain *Drosophila* tissues (Sturtevant et al. 1993; Karin 1994; Dickson 1995). In the *Drosophila* eye, *Drosophila* MAPK is capable of oppositely modulating the activities of two Ets-related factors, enhancing the Pointed activator (Brunner et al. 1994) and suppressing the Yan repressor (Rebay and Rubin 1995). In the latter case, MEK-activated MAPK phosphorylates Yan and thereby causes Yan to translocate back to the cytoplasm where it is rapidly subject to degradation, apparently elicited by carboxy-terminal PEST sequences (Rebay and Rubin 1995). We propose a similar explanation for the loss of CF2 where and when the kinase cascade is triggered by activation of EGF-R in the follicle cells. Interestingly, the CF2 amino acid sequence (Hsu et al. 1992) shows three PEST-like motifs (Rogers et al. 1986) in the middle region and, near the amino terminus, a putative site for phosphorylation by MAPK. We have observed that while CF2 disappears its RNA persists [data not shown], implying some post-transcriptional mechanism for CF2 clearance; we suggest a post-translational explanation, although we cannot exclude a translational mechanism.

Once initiated, the EGF-R cascade may operate as a loop, as it affects CF2 and CF2 affects rho, which in turn is known to affect but also to depend on EGF-R function (Ruohola-Baker et al. 1993). Loop function could amplify the dorsal signal (Sturtevant et al. 1993) and reinforce the assignment of dorsal cell fates. Initial transduction of the Grk signal occurs in the absence of Rho but may feed into the same pathway, initiating the loop [Fig. 7].

Our results indicate that CF2 functionally activates
ventral cell fates. We do not know whether it does so as a direct activator of ventral genes, as an indirect activator (e.g., through a double-repressor mechanism), or by default as a result of repression of dorsal genes. In any case, the influence of CF2 is propagated into the embryo, affecting the embryonic D/V pattern (Fig. 7).

Earlier Grk/EGF-R signaling is now known to specify posterior polar follicle cell fates, thus initiating anterior–posterior axis determination (Gonzales-Reyes et al. 1995; Roth et al. 1995). As in the case of Rho (Ruohola-Baker et al. 1993), CF2 does not seem to participate in this earlier patterning process; there is no evident suppression of CF2 in the posterior polar follicle cells (Fig. 1), and no anterior–posterior patterning defects have been observed upon CF2 misexpression (T. Hsu, unpubl.).

CF2 and subpopulations of the follicular epithelium

Follicular patterning involves not only the distinction between anterodorsal and ventral cells but the establishment of multiple, spatially distinct subpopulations of the follicular epithelium (Margaritis et al. 1980; Tolias and Kalafatos 1990; for review, see Spradling 1993). We note that the anterior-most border cells and the squamous cells associated with the nurse cell cap are devoid of CF2, in the wild type as well as in dorsalized and ventralized mutants (Fig. 1; data not shown); these cells do not express the AN296 enhancer trap marker (Fig. 2), despite the absence of CF2. Similarly, the cells that correspond to the two different elements of the rho expression pattern at stage 11 show different temporal susceptibilities to CF2. Evidently, unknown features of follicular cell differentiation delimit to specially competent subpopulations the ability to receive and respond to the EGF-R activation signal by loss of CF2 and expression of specific dorsal fates. At present, our results are consistent with CF2 having its major role at the earliest stage of D/V patterning, in a circumferential band of columnar follicle cells that surround the anterior portion of the oocyte. This band is divided in the wild type into CF2− and CF2+ sectors, as a result of EGF-R signaling; in dorsalized mutants the band is visualized by the circumferential lack of CF2 (Fig. 1). Future work using CF2 mutants and mitotic clone analysis might help illuminate the spatial domains and temporal dynamics of D/V patterning, and the roles of the CF2 transcription factor in specific aspects of this process.

Materials and methods

Drosophila strains

Genotypes of mutants (kindly provided by T. Schüpbach, Princeton University, NJ, and W.M. Gelbart, Harvard University, Cambridge, MA) used in this study were as follows: [1] pr, cn, topQ71, bw/CyO; [2] fs(1)K10^w, w, f/FM3; [3] b, pr, spiP48, bw/CyO; and [4] grkK830, cn, bw, E1p/CyO. AN296 is an enhancer trap/+/lacZ strain described by Bier et al. (1989), kindly provided by T. Schüpbach. The fly strains fs(1)K10^w, w, f/FM3; P[w+, hsp70-CF2] (K10/CF2-s for short) and w, pr, cn, topQ71, bw/S66a; P[w+, hsp70-CF2] (top/S66; CF2-s for short) were obtained by standard crosses.

Transgenic lines

For CF2 overexpression experiments, CF2-s transgenic lines were created using a 2-kb Eagl-ClaI fragment of the ovarian CF2 cDNA (Gogos et al. 1992, Hsu et al. 1992) containing 119 bp of the 5′-untranslated region [UTR], the entire coding region, and 429 bp of the 3′ UTR. This fragment was cloned into the Hpa1 site of the pCaSpeR-hs vector (Pirrotta 1988) in the forward orientation downstream of the hsp70 gene promoter. For CF2 depletion experiments CF2-as lines were created using an 887-bp HindIII (from the plasmid vector)–NarI fragment derived from a partial ovarian CF2 cDNA clone (Gogos et al. 1992, Hsu et al. 1992), containing 247 bp of the 5′ UTR and 640 bp of the 5′ portion of the coding region. This fragment was cloned into the Hpa1 site of the pCaSpeR–hs vector in the reversed orientation. Germ-line transformations in the yw strain were performed by standard procedures (Spradling 1986). Five CF2-s homozygous lines were generated that exhibited similar phenotypes upon heat treatment; the data reported came from line T3, bearing the insert on the third chromosome. Three CF2-as homozygous lines were obtained and crossed to generate two doubly homozygous lines; we show data from line as1,aT1, with inserts in both autosomes.

Heat treatment

Newly hatched females were conditioned (with half the number of males) in the presence of live yeast at room temperature for 4 days. Three types of heat shock regimens were used: (1) low level: 3-day incubation at 30°C; (2) high level: 3-day incubation at 30°C plus two 15-min incubations at 37°C daily (e.g., at 9 a.m. and 7 p.m.); (3) pulsed heat shock: flies were transferred to a prewarmed bottle containing media and incubated at 38–39°C for 30 min, after which the bottles were transferred to 25°C and incubated for 30 min. We determined that if no further disturbance (such as anesthesia) was applied at this point, the flies would resume egg laying. The entire heat shock and recovery regimen is referred to as heat treatment. After heat treatment, the flies were either transferred to fresh bottles of media, incubated at 25°C, and dissected at different times for examination of ovaries, or were transferred to cages for egg collection and scoring of embryos as described (Roberts 1986).

Mosaic analysis

The strain Df(2L)ry7^1 cn/CyO; ry50k, a kind gift of J. Tower (University of Southern California, Los Angeles) was generated by imprecise P-element excision after y-irradiation of the homozygous viable starter strain P[acman], ryY^-P[63S, cn. The original insertion was mapped 5–10 kb upstream of CF2, and the resulting deletion removes ~25 kb [J. Tower, pers. comm.]. The deletion results in a larval lethal phenotype and was characterized by in situ hybridization of CF2 genomic clones to polytene chromosomes, genomic PCR, and CF2 antibody staining of homozygous embryos; no CF2 coding region or CF2 protein expression could be detected [C. Baggi, S. Bolshakov, and C. Mollinari, unpubl.]. Female yw/+, Df(2L)ry7^1 cn/+ and control yw homozygotes were X-ray-irradiated (1050 R, Torex 120D source) as first-instar larvae or adults according to the method of Stevens et al. (1990). Eggs were collected on apple juice/agar plates for a period of 3 days, and dorsal appendages were examined. Selected eggs were either critical-point dried or air-dried before sputter-coating and viewed using a Philips XL-30 field emission scanning electron microscope. Images were processed using Adobe Photoshop.

Immunostaining

The CF2 monoclonal antibody was obtained from a mouse hy-
Hsu et al.

formaldehyde (from Polyscience) in PBS, washed, and permeabilized for 8 min in PBS containing 0.1% Triton X-100. After three washes in PBS, follicles were incubated with CF2 monoclonal antibody for 3–15 hr, washed three times in PBS, and incubated for 1–2 hr with 1:300 dilution of fluorescein |DTAF)-conjugated affinity pure goat anti-mouse IgG [H + L] (from Jackson ImmunoResearch Laboratories) in PBS, 0.5% BSA. They were rinsed for 10 min in PBS and transferred through a series of 20-min incubations in PBS solutions with increasing concentrations of glycerol: 30%, 50%, and 80% (adjusted to pH 7). Whole mounts were prepared by transferring follicles into Gelvatol mounting solution [Harlow and Lane 1988]. They were analyzed using a Compact Confocal Microscope (CCM) built at the European Molecular Biology Laboratory (EMBL), Heidelberg, Germany [E. Stelzer and collaborators, Light Microscopy Group].

Detection of β-galactosidase activity in ovaries

Ovaries bearing the AN296 enhancer trap insert were dissected in PBS from well-conditioned and pulse heat-shocked females (see above) at 1.5 hr after heat treatment and were fixed for 8 min in 1% glutaraldehyde (from Sigma) in PBS. They were rinsed three times in PBS 0.5% Triton-X100, stained at 37°C for 30 min in PBS containing 0.2% X-gal (from Sigma), 5 mM K4[Fe(III)]CN6, 5 mM K3[Fe(II)CN6], 0.3% Triton-X 100, rinsed in PBS, and mounted in PBS 90% glycerol for microscopic analysis.

RNA in situ hybridization

The hybridization probes used were a 2.5-kb rho cDNA fragment encompassing the entire 1.07-kb coding region (kindly provided by H. Ruohola-Baker, University of Washington, Seattle), a 4.5-kb dpp cDNA from the pNB40 library (St. Johnston et al. 1990), and a 530-bp twi genomic EcoRI-BamHI fragment (Thiese et al. 1988), the dpp- and twi-containing plasmids were kindly provided by W. Gelbart). The probes were labeled with digoxigenin as described by the supplier [Boehringer Mannheim Biochemicals]. For rho staining of egg chambers, newly hatched females were conditioned and pulse heat-shocked as described above, kept in the same bottle of media for another 1.5 hr of incubation at 25°C, and anesthetized on ice; ovaries were dissected in Ringer's solution. For dpp and twi staining of embryos, well-conditioned young flies (4–5 days old) were heat-shocked at the high level as described, and embryos were collected at 14–18 hr after heat treatment and incubated for another hour at 25°C (i.e., collection of 1–5 hr embryos). The staining and detection procedures have been described [Tautz and Pfeifle 1989, Hsu et al. 1993]. For examination, ovaries and embryos were mounted in a mixture of Ringer's and glycerol [1:1] or a mixture of PBS/0.1% Tween 20 and glycerol [1:1], respectively.

Cuticle preparations

Cuticle preparations were made essentially as described [Wiechaus and Nüsslein-Volhard 1986], and photographed by dark-field optics.

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