In vitro transcription of the *Drosophila engrailed* gene

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An enzyme system that accurately initiates transcription of the *engrailed* gene has been prepared from *Drosophila* embryos. The system has been separated chromatographically into two fractions, both of which are required for specific *engrailed* transcription. DNase footprint and competition analysis detected at least two sequence-specific DNA-binding proteins in one of these two fractions. Together, these proteins bind to eight regions within 400 bp of the transcription initiation sites. Most of the regions containing these binding sites are required for minimal *engrailed* transcription in vitro. In addition, a region downstream from the initiation sites and within the first 40 residues of the transcription unit is essential for transcription. Transient in vivo expression assays indicated that these same upstream and downstream sequences are required for transcription in *Drosophila* tissue culture cells.

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The structure of the *Drosophila engrailed* gene is highly unusual. Mutations in the gene are spread over a 70 kb region of the genome, yet its single transcription unit is less than 4 kb in length [Drees et al. 1987]. Flanking its transcription unit are transcriptionally silent regions that extend for 20 kb downstream and 45 kb upstream, suggesting complex and perhaps novel regulatory mechanisms. To understand the role of these flanking regions, and learn how expression of the *engrailed* gene is regulated, we have developed a system that faithfully initiates *engrailed* transcripts in vitro.

The central importance of transcriptional regulation in generating the temporal and spatial patterns of *engrailed* gene expression has been suggested by several observations. Mutations that alter the structure of the gene's untranscribed flanking regions affect both the quantity and disposition of *engrailed* transcripts [Weir and Kornberg 1985; and unpubl.]. Also, the distinctive expression in the early embryo, in which single-cell-wide stripes of cells express the *engrailed* gene in each of the segmental primordia, is the result of an extremely rapid stepwise progression in a temporal pathway [Weir and Kornberg 1985]. This progression initiates with 3 broad stripes, proceeds to a pattern with multiple stripes that alternate in intensity, and eventually results in 14 equally spaced zebra-like bands in blastoderm stage embryos [T. Karr, M. Weir, and T. Kornberg, in prep.]. At this developmental stage, *engrailed* RNA has a half-life of approximately 15 min [M. Weir and T. Kornberg, unpubl.], and the progression of patterns is completed within 90 min. Such rapid and dramatic changes are likely to be under transcriptional control. Regulation of *engrailed* gene expression has been investigated genetically. The *engrailed* gene is not expressed in *even-skipped* mutant embryos [Harding et al. 1986], and *engrailed* gene expression is also affected in *hairy* and *fushi tarazu* mutant embryos [Howard and Ingham 1986]. Thus, the *even-skipped*, *hairy*, and *fushi tarazu* genes are required for the normal expression patterns of the *engrailed* gene. Although these studies do not indicate how this requirement is manifested at a molecular level, it is possible that the protein products of these genes directly regulate *engrailed* gene transcription.

An alternate approach to probing the mechanisms that control expression of the *engrailed* gene is to reconstruct the transcriptional regulation in vitro. In vitro systems from many cell and animal sources have been described that accurately reconstruct RNA polymerase II initiation [Dignam et al. 1983; Manley 1983]. Several systems derived from mammalian sources have been shown to exhibit regulation in response to tissue-specific factors [Mizushima-Sugano and Roeder 1986; Gorski et al. 1986]. In *Drosophila* systems, extracts of *Drosophila* tissue culture cell nuclei have been shown to initiate accurately from the promoters of *Drosophila* heat shock, alcohol dehydrogenase, actin, and histone genes [Parker and Topol 1984a,b; Heberlein et al. 1985]. Such tissue-culture-derived systems unfortunately yield only small amounts of enzymes for biochemical characterization, and they cannot take advantage of either the rich genetic heritage of *Drosophila* or of its readily accessible developmentally staged populations. In this study, we have developed a method to prepare extracts from *Drosophila*.
embryos, and have shown that these extracts can accurately initiate transcription of several *Drosophila* genes. We use this system to investigate transcription of the *engrailed* gene with the aim of elucidating the complex mechanisms that regulate this transcription.

**Results**

**In vitro transcription in *Drosophila* embryo extracts**

A cell-free extract of *Drosophila* embryos was prepared and assayed for its ability to transcribe a number of templates containing cloned *Drosophila* promoters. These templates were cleaved at sites downstream from the in vivo initiation sites and were used in run-off assays (Fig. 1A). The major run-off RNA product, 1.1 kb, 1.2 kb, and 1.4 kb in length for the *fushi tarazu*, alcohol dehydrogenase (*Adh*), and actin 5C genes, respectively, in each case corresponds to a transcript initiating at the in vivo start sites (Fyrberg et al. 1980; Benyajati et al. 1983; Laughon and Scott 1984). The size of the run-off transcript from an *engrailed* template (1.3 kb) was also consistent with the in vivo initiation sites (Drees et al. 1987). Under these conditions of template and extract concentrations, the relative efficiencies of the *fushi tarazu*, *Adh*, actin 5C, and *engrailed* templates were approximately 10:5:4:1, respectively. The *fushi tarazu* DNA template, the rate of product accumulation was constant for a 30-min incubation (data not shown), and the amount of RNA synthesized was proportional to the template concentration. Maximal incorporation was at 40 µg/ml DNA in the presence of 125 µg crude extract (Fig. 1B).

A primer extension assay was used to compare the in vitro initiation sites with the sites utilized in vivo. A 5'-end-labeled oligonucleotide was hybridized to RNA that had been synthesized in vitro or isolated from *Drosophila* embryos, these were elongated with reverse transcriptase, and the products were fractionated by electrophoresis (Fig. 2A). Transcription initiates in vitro at a cluster of closely spaced sites identical to those used in vivo. These are designated by their nucleotide spacing in the direction of transcription, with the major sites of initiation at residues +1, +2, +5, and +6. The *Drosophila* embryo extracts therefore accurately initiate transcription from the *engrailed* promoter.

**Fractionation of the embryo extract**

The embryo extract was fractionated by a procedure that has been used previously to identify transcription components of HeLa cells (Dynan and Tjian 1983) and *Drosophila* tissue culture Kc cells (Heberlein et al. 1985). After dialysis, the crude extract was applied to a heparin-Sepharose column and eluted with 0.4 M KCl (fraction H.4; nomenclature of Heberlein et al. 1985). The H.4 fraction was applied to a DEAE-Sepharose column from which a break-through fraction (D.1) and a

![Figure 1](image-url)
fraction that contained proteins that eluted with 0.25 M KCl (D.25 fraction) were collected.

Each of these protein fractions was tested for transcriptional activity on an engrailed template using a primer extension assay (Fig. 3). The crude extract and H.4 fractions alone were active, whereas the D.1 and D.25 fractions were not. However, transcription activity was restored when the D.1 and D.25 fractions were combined. Transcription activity was sensitive to 0.5 μg/ml α-amanitin, indicating that RNA polymerase II is an essential component. Although the identities of the other components of these DEAE fractions have yet to be established, RNA polymerase II activity was unevenly distributed between them. On a template of single-stranded calf thymus DNA, 90% of the RNA polymerase II activity was present in the D.25 fraction. These results suggest that both DEAE fractions contain essential components: the D.25 has RNA polymerase II activity and the D.1 has some component(s) that enhance(s) specific transcriptional initiation from the engrailed promoter. Since the D.25 fraction could not be replaced in the reaction by a preparation of RNA polymerase II purified from Drosophila embryos (Coulter and Greenleaf 1982), either a distinct RNA polymerase II subspecies in the D.25 fraction is involved, or the RNA polymerase purified by an assay using a nonselective calf thymus DNA template is inactive in this initiation reaction (Kim and Dahmus 1986), or the required D.25 component(s) include additional activities.
**D.1 D.25**

**A** -52E

**B** -85

**C** -105

**D** -138

**E** -152

**F** -181

**G** -197

**H** -230

**I** -250

**J** -263

**K** -283

**L** -38

**M** -241

**N** -380

**O** -419

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**Figure 4.** Footprint analysis of the *engrailed* promoter upstream of its initiation sites. Reactions were carried out as described in Experimental procedures. The probe extended from position +231 to −305 and was labeled at the −305 position by a 5′-end-filling reaction. Shown is the region from +20 to −300. (Lanes 1 and 4) DNase I digests of the probe in the absence of any embryo protein fraction. (Lanes 2 and 3) Binding reactions and DNase digests in the presence of 30 and 60 μg fraction D.1 protein, respectively. (Lanes 5 and 6) Reactions were carried out in the presence of 15 and 30 μg fraction D.25 protein. Regions of reduced or absent DNase cleavage sites are bracketed and labeled A–G. The upturned arrow marks the position of the first of the four initiation sites. Dashed brackets refer to regions of weak protection from DNase cleavage. The unlabeled bracket adjacent to lane 6 denotes the target sequence of a new binding activity present in the D.25 fraction and described in the text.

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**DNase I footprint analysis of the engrailed promoter**

To analyze the activity conferring specificity on initiation of *engrailed* transcription, the capacity of the D.1 and D.25 fractions to protect the promoter region from DNase I digestion was determined. Several probes were used. The first was designed to investigate the first several hundred nucleotides of sequence upstream of the start sites. It extended from position −305 to +231 and was labeled at position −305. Increasing amounts of D.1 were incubated with this probe prior to partial degradation with DNase I [Fig. 4]. Seven distinct regions of protection were detected (labeled A–G), the most proximal site (A) mapping to position −34 to −52, and the most distal site (G) mapping between positions −263 and −283. The lengths of the protected regions varied from 14 to 22 residues, and the distances between any two adjacent sites (other than B and C) varied from 11 to 38 residues. The extent of cleavage in the region between sites A and B was strongly enhanced in the presence of some preparations of the D.1 (or H.4, not shown) protein fractions [Fig. 4] but not others [Fig. 5]. The source of this variability is not known.

In contrast, D.25 retained little sequence-specific binding activity. With D.25, regions E and F were only partially protected, and no protection was observed for the other regions. This experiment did reveal the presence of a new binding activity whose target sequence (−9 to +10) includes the initiation sites. This activity could represent a subspecies of RNA polymerase II, or it could be one of its accessory factors. Purified RNA polymerase II did not bind to this region in these footprint reactions [data not shown].

Using a DNA fragment that extended further upstream, from residues −38 to −419, a protected sequence, called region H, was found to extend from nucleotides −305 to −348 [Fig. 5A]. The length of this region is more than twice that of any of the seven other binding sites, and it contains four copies of the pentamer sequence GAGAG (CTCTC). This sequence is also in the center of regions A, B, C, E, and F. Of the protected regions in which this sequence was not found, region D has repeats of the sequence GAGTG (or CACTC). Re-
Figure 5. (See facing page for legend.)
region G, which has imperfect dyad symmetry, has no similar sequence. The nucleotide sequence from -422 to +216 and the footprinted regions are shown in Figure 5C.

The number of distinct binding activities responsible for the eight protected regions was established by competition binding assays. A double-stranded oligonucleotide consisting of the 19 residues of region C was incubated in the presence of the -38/-419 probe and fraction H.4 (Fig. 5A). As the ratio of unlabeled competitor DNA to probe DNA was increased, protection at the six binding sites that contain the GAGAG pentamer disappeared. Region D contains the related pentanucleotide GAGTG, and its protection was consistently weak relative to the other regions, suggesting that its affinity for the binding protein is weak. The footprint of region D was also competed by the region C oligonucleotide. Only region G protection was unaffected in the presence of the region C competitor. In contrast, when a 60-bp AvaI/Ddel DNA fragment containing region G was used as a competitor, it suppressed protection only at region G (Fig. 5B). Thus, there appear to be two distinct binding activities specific for the engrailed promoter: one binds to sites containing the pentanucleotide sequence GAGAG (or CTCTC); the other recognizes a single site, region G, whose sequence contains dyad symmetry.

**Figure 5.** Footprint analysis of the engrailed promoter in the presence of competitor DNAs. (A) A probe extending from position -38 to -419 was labeled at either the downstream [coding] end or the downstream noncoding end by a 3'-end-filling reaction. Size markers were from Maxam and Gilbert G/A sequencing reactions. DNase digestions were carried out as described in Experimental procedures with the following additions: (lanes 1 and 8) no added protein fraction; (lanes 2 and 9) 1.4 ~g fraction H.4; (lanes 3 and 4) same as lane 2 except that a 4- and 40-fold molar excess of sites, respectively; see text). (B) Same fragment and procedure as in A with only the noncoding strand labeled with polynucleotide kinase: (lanes 1 and 8) no added extract protein; (lane 2) 14 ~g fraction H.4 and 1.0 ~g poly(dI/dC) (as a nonspecific competitor); (lanes 3-7, 12-14) identical to lanes 4 and 11 except for increasing amounts of region C oligonucleotide as competitor [4, 40, and 400-fold molar excess of sites, respectively; see text]. (C) The DNA sequence of the coding strand surrounding the engrailed initiation sites is shown and the location of the eight footprinted regions on the engrailed promoter are indicated. The four initiation sites are represented by > and the first is labeled as + 1. The engrailed translational initiation codon is presented in larger and bold type.

**Upstream sequences are required for engrailed transcription**

If the DNA-binding proteins that recognize sequences upstream of the engrailed initiation sites are involved in promoting transcription, then removing these upstream sequences should affect the fidelity or efficiency of RNA polymerase II initiation. To evaluate the role of these sequences, 18 templates were constructed that retained various lengths of engrailed DNA upstream of the initiation sites [Fig. 6A]. These templates ranged from Δ - 887 (retaining 887 bp upstream of the first initiation site and 231 bp downstream) to Δ + 15 (retaining only engrailed DNA from the downstream positions +15 to +231). Relative transcription efficiencies of these DNAs were determined. Each reaction mixture also contained a control template consisting of engrailed promoter DNA with a 15-bp insertion downstream from the initiation sites. Transcripts from this internal control template could be readily distinguished by their increased length, permitting a direct comparison of the template efficiencies of the different promoter deletions. Template efficiencies of 18 promoter fragments were compared qualitatively by inspection of autoradiograms after electrophoretic separation of their primer extension products. Quantitative comparisons were made from densitometric scans of the autoradiograms. Densitometric scans of the autoradiograms.
metric values were compared to that of template Δ - 305, which was arbitrarily assigned a value of 100% (Fig. 6A).

A direct correlation between promoter length and template efficiency was observed: As the length of upstream sequence decreased, transcriptional activity decreased in discrete steps. Activity decreased twofold with template Δ - 130, decreased little further with templates Δ - 118, Δ - 112, and Δ - 101, and decreased an additional threefold with template Δ - 67. These quantitative changes correlate with the presence of protected regions B and C between positions -67 and -105, and regions D through G between positions -138 and -283. No change in the relative yield from the four start sites was observed. These results suggest that the sequence-specific DNA-binding activities specific for the engrailed promoter play a major role in the efficiency of the initiation reaction.

A similar conclusion was suggested by the behavior of mutant templates with an internal deletion. These deletion mutants lack between 16 and 49 bp within the region bounded by positions -80 and -26, and are designated by the borders of their deletion end points. Templates Δ -80/-38, and Δ -77/-26 remove sequences that include both regions A and B, their transcription was reduced by approximately 85%. Templates Δ -51/-35 and Δ -65/-34 remove only sequences within and/or around region A, their activities were reduced by 83-91%. The fact that deletion of region A alone affects transcription as extensively as deletion of...
region A and B argues either that spacing of region A relative to the initiation sites affects template efficiency or that this most proximal binding site plays a special role in this promoter. However, these results are still consistent with the hypothesis that the sequence-specific DNA-binding activities detected in this region are intimately involved with *en grailed* transcription.

All templates that retained 14 or fewer residues upstream of position +1 supported initiation only from sites +5 and +6. The inability of the Δ−14 template or the Δ−77/−26 internal deletion mutant to utilize initiation sites +1 and +2 suggests that *en grailed* DNA sequences between position −14 and −34 (the most proximal residue of DNA-binding region A) are used by the transcription apparatus in scanning the promoter sequences for the normal initiation sites. Interestingly, transcription was detectable from template Δ+4, even though all upstream sequences, including initiation sites +1 and +2, are deleted, and only a single base pair upstream of initiation site +5 is retained. The capacity of templates lacking all of the upstream promoter sequences to support transcription, albeit at a low level, implies that sequences within the transcribed region are recognized by the transcriptional machinery that initiates RNA synthesis.

**A region within the en grailed transcription unit is essential for transcription in vitro**

To determine if sequences within the nontranslated leader of the *en grailed* transcription unit are involved in the initiation reaction, templates were constructed that deleted downstream sequences. These leader sequence deletion mutants had varying amounts of DNA deleted between positions −7 and +127, but all retained in common sequences upstream from −7 and downstream from +127 [Fig. 6B; designation of mutants is by their deletion endpoint closest to residue +1]. Transcriptional activity of these DNAs was measured using a primer extension assay in the presence of the control insertion template and the crude extract. Since these deletion mutants remove sequences from within the transcription unit, the lengths of the transcripts and of the primer extension products were correspondingly shortened. Removal of sequences between +39 and +124 did not affect initiation quantitatively or qualitatively. In contrast, the transcription efficiency of the Δ+37 DNA (which lacks sequences between +37 and +124) was lowered at all four sites by approximately 75%. The Δ+30 mutant was essentially inactive. We conclude that there is a region within the first 38 residues downstream of site +1 that is essential for transcription.

**Region C competitor DNA specifically blocks high level transcription of the en grailed promoter**

To further evaluate the role of the DNA-binding proteins in promoting transcription, the activities of templates Δ−118 and Δ−34 were determined in the presence of region C oligonucleotide at concentrations previously shown [Fig. 5A] to suppress protection from DNase I digestion in regions A through F and region H. In these experiments, the Adh proximal promoter served as the control template since this promoter does not contain the GAGAG (or CTCTC) pentamer. Again, template efficiencies were determined from densitometric scans of the autoradiograms [Fig. 7]. Transcription of template Δ−118, which contains regions A, B, and C, decreased to 22% of its maximal level as the ratio of competitor DNA to template was increased. In contrast, the competitor had no appreciable effect on transcription of template Δ−34. This template has all upstream binding sites deleted; its activity in the absence of competitor is comparable to template Δ−118 in the presence of competitor. These results confirm the notion that the proteins that bind to the GAGAG (or CTCTC) pentamer sites play a major role in modulating this gene’s transcription.

Similar experiments were carried out using the 60-bp *AvaI/DdeI* DNA fragment containing region G as the competitor [data not shown]. No decrease in transcription was detected with templates containing region G. Thus, it is possible that the DNA-binding activity that recognizes region G plays no role in *en grailed* transcription per se. However, given the sizable distance (263 bp) of region G from the initiation sites, the failure to detect an effect on transcription is more likely due to the insensitivity of in vitro transcription assays to far upstream sequences. Efforts to address this question by moving the region G sequence closer to the initiation sites are currently under way.

**In vivo transcription has similar upstream and downstream sequence requirements**

To evaluate the importance of the sequences implicated by in vitro initiation assays, transcription was also quantitated in *Drosophila* tissue culture cells. Various lengths of upstream and downstream promoter sequences were fused with sequences encoding the enzyme chloramphenicol acetyl transferase (CAT), and plasmids with these fusion gene constructs were transfected into *Drosophila* Schneider 2 cells. To control for transfection efficiency, a plasmid containing a fusion of approximately 3.5 kb of the upstream *en grailed* promoter with the coding sequences of *E. coli lac Z* was transfected simultaneously. Forty-eight hours after transfection, extracts from the transfected cells were prepared, and CAT and β-galactosidase activities were measured.

CAT fusion genes with various lengths of upstream promoter sequences retained the first 127 bp downstream from the first initiation site. As with the in vitro primer extension experiments, template Δ−365DM26 was arbitrarily assigned a transcriptional value of 100% [Fig. 8A]. CAT activity decreased 3-fold with template Δ−118DM26, and 11-fold with template Δ−38DM26. Template Δ+2DM26 reproducibly retained only 3% of the maximal transcription activity.

Four CAT fusion genes with various lengths of the
nontranslated leader were examined. These templates retained 118 bp of upstream sequence, and either 127, 39, 37, or 30 bp of downstream sequence. Template Δ+127DM26 was assigned a transcriptional value of 100% (Fig. 8B). CAT activity in extracts of cells transfected with template Δ+39DM26 was reduced by 54% (Compared to a 10% decrease that was observed in vitro, see Fig. 6B). The relative activity of template Δ+37DM26 (23%), was virtually identical to the in vitro efficiency. Template Δ+30DM26 had only 3% activity relative to template Δ+127DM26, thus confirming that sequences within the transcribed region of the engrailed gene are important for transcription. We conclude that both in vivo transcription activity and in vitro initiation efficiency depend on the same upstream and downstream sequences, and that the in vitro assays portray the initiation reaction faithfully.

Discussion

In this first investigation into the molecular mechanisms that directly regulate expression of the engrailed gene, we found that the engrailed promoter is unusual in several respects. It has neither a TATA box nor a CAAT element, sequences found in many eukaryotic promoters and thought to be important for RNA polymerase II recognition [Shenk 1981]. Perhaps the absence of these sequences accounts for the multiple sites of initiation we observed for the engrailed gene in vitro and in vivo, and for the relatively low activity of the engrailed promoter in the in vitro transcription reaction (Fig. 1). The other promoters assayed in the in vitro system [Adh, Actin 5C, and fushi tarazu] have TATA homologies and are all approximately an order of magnitude more efficient. This suggests that some sequence and protein components that confer specificity to RNA polymerase are different for the engrailed gene than for these other genes.

Features of the engrailed promoter that were unexpected in view of the extraordinary size of the engrailed regulatory region were the high density of functional sequence elements within the first 400 bp upstream of the sites of initiation, and the presence of a critical downstream sequence element. Eight distinct regions in this upstream region were recognized by sequence-specific DNA-binding activities. Interestingly, competition DNA-binding studies suggested that seven of the eight binding sites are recognized by the same protein, implying that several copies of this protein are involved in initiating engrailed transcription. Since binding was observed in the absence of RNA polymerase II, the function of the DNA binding is likely to be in the early steps of the initiation process [Hawley and Roeder 1985].

The identified downstream sequence is necessary and, for two of the four sites of initiation, sufficient for at least low levels of transcription. Deletion mutants that lack all of the upstream sequences and retain only the residues four nucleotides downstream from the first initiation site are nevertheless active for the two remaining sites of initiation. The overall template efficiency of such mutants was reduced 10-fold relative to templates with 300 upstream residues. The nature of the role of downstream sequences is unclear, although the close proximity of the essential region to the initiation sites suggests that the RNA polymerase or proteins with which it is directly associated interact with the sequence. A similar requirement for downstream sequences has been observed previously for the Drosophila
Ultrabithorax (M. Biggin, pers. comm.) and 74E ecdysone-responsive (C. Thummel, pers. comm.) genes.

Although we have not yet identified the proteins that bind to sequences within 350 bp upstream of the sites of initiation, it is unlikely that they are previously isolated transcription factors. For example, no detectable homology was found between these regions of the *engrailed* promoter and the sequences recognized by either HSTF, a heat shock factor (Parker and Topol 1984b), or Adf1, an Adh gene binding protein (Heberlein et al. 1985). Several aspects of the binding sequences within the *engrailed* promoter are, however, noteworthy.

Six of the eight upstream regions (regions A, B, C, E, F, and H) have in common the sequence GAGAG in the middle of each site, but little additional homology. Region D contains the related sequence GAGTG. Given that competition studies suggested that these sites are recognized by the same binding activity, the pentamer may form a core recognition sequence for the same DNA-binding protein. The importance of region A is also suggested by the existence of a homologous sequence in another *Drosophila* gene. Eleven of thirteen residues (−39 to −52) in region A are homologous with a sequence in the *invected* gene (K. Coleman and T. Kornberg, unpubl.). The region of homology in the *invected* sequence is in a similar location relative to the site of transcription initiation (from positions −23 to −37). The *invected* sequence is also protected from DNase I cleavage by DNA-binding activities in the H.4 and D.1 fractions (W. Soeller, K. Coleman, and T. Kornberg, unpubl.), as is region A of *engrailed*. Since the *invected* gene is closely related to the *engrailed* gene in sequence and pattern of expression (Poole et al. 1985) and it is also expressed exclusively in posterior compartment cells (Coleman et al. 1987), we speculate that homologous sequences in the promoter regions of these genes help to orchestrate their coordinate regulation.

We have noted other homologies in the regions protected from DNase I digestion. Region B contains 9 of 13 base pairs identical to an upstream region of the *Ultra-*

![Figure 8. Transient expression assays of upstream and downstream deletion mutant templates. (A) *engrailed* templates labeled Δ-DM26 and containing various lengths of upstream sequence were fused to a reporter gene coding for CAT as described in Experimental procedures and were transfected with a β-galactosidase control template into Schneider 2 cells. (% conversion) The fraction of chloramphenicol substrate converted to acetylated forms. DM26 is the CAT plasmid vector with no inserted promoter present. Results reflect values obtained after normalizing β-galactosidase activity measured in each sample and subtracting background levels determined by transfection with pDM26 alone. CAT assays were carried out as described by Gorman et al. (1982) at 37°C for 5 min using 50 μg protein extract per assay. (B) Corresponding *engrailed* templates with downstream deletions fused to the CAT reporter gene.](https://genesdev.cshlp.org/article/8/1/165/107941)
bithorax promoter that can be protected from DNase I digestion [M. Biggin, pers. comm.]. Likewise, region C has a significant degree of identity with an additional upstream protected region of the Ultrabithorax gene (M. Biggin, pers. comm.), and an upstream region of the fushi tarazu gene (~102 to ~114 relative to the start site reported by Laughon and Scott 1984). Region G shares no obvious homology with the other six engrailed upstream sites. However, it does possess a significant degree of dyad symmetry. Such symmetry in a DNA-binding site is characteristic of DNA sequences bound by certain transcription factors [Topol et al. 1985; Ptashne 1986]. In addition, this region shares a high degree of sequence similarity with an element discovered in the dopa decarboxylase [Ddc] promoter of Drosophila melanogaster and D. virilis. This element, situated between positions ~56 and ~73 in the D. melanogaster promoter, is identical to engrailed region G at 10 of 13 nucleotide residues (overlapping but not entirely coincident with the stretch of dyad symmetry). It has been shown to be essential for expression of Ddc in both the larval and adult nervous system [Schonlick et al. 1986]. Since engrailed function is required in the adult nervous system (Lawrence and Johnston 1984) and since engrailed is expressed in the embryonic nervous system [DiNardo et al. 1985; C. Hama and T. Kornberg, in prep.], one may speculate that region G and its binding protein are involved in central nervous system-specific expression. This notion is consistent with our preliminary results that indicate the region G binding activity to be present in embryo but not tissue culture extracts [data not shown].

We have demonstrated that the activity responsible for protection of the GAGAG sites from DNase digestion also modulates maximal engrailed transcription in vitro. Whether this activity or the GAGAG sites are involved in conferring spatial or stage specificity on engrailed expression is not known. However, results of P-element-mediated transformation experiments utilizing engrailed promoter-β-galactosidase DNA fusion constructions argue that these sites are not sufficient for tissue or stage specificity. Transformants containing constructs with only 1 kb of upstream sequence do not display a clear stripping pattern. For a wild-type engrailed expression pattern to develop in the epidermis and central nervous system of the embryo, at least 7.5 kb of upstream sequence are required [C. Hama and T. Kornberg, in prep.]. Dissection and mapping of cis-acting elements within this 7.5-kb region by such transformation experiments will allow us to test promising candidates for their effect on engrailed transcription in vitro. Such an approach should allow us to identify, purify, and characterize trans-acting activities responsible for the normal tissue-specific gene regulation.

It will be interesting to determine whether some of the activities responsible for protecting the engrailed promoter from DNase I digestion are also responsible for modulating transcription of the Ultrabithorax, fushi tarazu, even-skipped, and engrailed have cross-regulatory interactions. In vitro transcription reactions should help us understand the nature of these interactions.

**Experimental procedures**

**Preparation of the embryo extract**

Oregon R wild-type flies were grown under standard conditions at 25°C in population cages [Kriegstein and Hogness 1974]. Embryos of various ages, usually 2–12 hr after egg laying, were collected on corn meal–agar trays, harvested with a solution of 0.7% NaCl, 0.04% Triton X-100 (Sigma), and dechorionated by immersion in 2.25% sodium hypochlorite (1/2 dilution of commercial bleach) for 90 sec. After extensive rinsing with sodium chloride/Triton solution and a final rinse with H2O, the embryos were quickly blotted dry and weighed. All subsequent steps were carried out at 4°C.

The initial steps for isolation of nuclei closely follows the procedure described in Kuo et al. 1982. The embryos were suspended in 4 ml/gram of embryos in buffer A [0.35 M sucrose, 15 mM HEPES [pH 7.6], 10 mM KCl, 5 mM MgCl2, 0.1 mM EDTA, 0.5 mM EGTA, 1 mM DTT, 1 mM PMSF, 1 μg/ml leupeptin [Boehringer-Mannheim]], and were homogenized by six strokes with a motorized Teflon pestle (G.K. Heller, Floral Park, New York) followed by six strokes with a glass–glass dounce homogenizer [type B pestle]. The milky-white homogenate was passed through three layers of pre-wetted Miracloth [Calbiochem] and centrifuged at 4000 rpm in a Beckman JA20 rotor for 10 min. Pellets were resuspended gently in buffer A (2 ml/g embryos), and the nuclei in 6-ml aliquots were layered over an equal volume of buffer B [0.8 M sucrose, 15 mM HEPES [pH 7.6], 10 mM KCl, 5 mM MgCl2, 0.1 mM EDTA, 1 mM EGTA, 1 mM DTT, 1 mM PMSF, 10 μg/ml leupeptin] and centrifuged 4000 rpm in a Beckman JS-14 rotor at 5000 rpm for 10 min. Subsequent steps closely followed the procedure for preparation of K, nuclear extracts [Parker and Topol 1984a; Heberlein et al. 1985]. After noting the volume of the nuclear pellet and removing the supernatant by aspiration, nuclei were resuspended in 4 volumes of buffer C [15 mM HEPES [pH 7.6], 110 mM KCl, 3 mM MgCl2, 0.1 mM EDTA, 1 mM DTT, 1 mM PMSF, 10 μg/ml leupeptin] and transferred to Beckmann 60Ti polycarbonate tubes. Nuclei were pelleted by centrifugation in a Beckman JS-14 rotor at 5000 rpm for 10 min.

Subsequent steps closely followed the procedure for preparation of K, nuclear extracts [Parker and Topol 1984a; Heberlein et al. 1985]. After noting the volume of the nuclear pellet and removing the supernatant by aspiration, nuclei were resuspended in 4 volumes of buffer C [15 mM HEPES [pH 7.6], 110 mM KCl, 3 mM MgCl2, 0.1 mM EDTA, 1 mM DTT, 1 mM PMSF, 10 μg/ml leupeptin] and transferred to Beckmann 60Ti polycarbonate tubes. Ammonium sulfate 4.0 M, [pH 7.6] was added stepwise to a final concentration of 0.36 mM, and lysis of the nuclei was accomplished by gentle mixing on a nutator [Clay Adams] for 30 min. The extract was clarified by centrifugation at 35,000 rpm for 1 hr in a Beckman 60Ti rotor. Material floating on the supernatant surface was removed by aspiration. The supernatant was collected by aspiration, and the active protein fraction was precipitated by the stepwise addition of 0.3 grams of solid ammonium sulfate per milliliter of supernatant. After stirring for 15 min, the precipitate was collected by centrifugation at 15,000 rpm for 15 min in a JA20 rotor. The pellet was gently resuspended with a glass rod in a volume of buffer HEMG [25 mM HEPES [pH 7.6], 40 mM KCl, 12.5 mM MgCl2, 0.1 mM EDTA, 1 mM DTT, 1 mM PMSF, 10 μg/ml leupeptin, 10% glycerol [see Heberlein et al. 1985] equal to one-third the volume of the nuclear pellet. The resulting suspension was clarified by centrifugation [10,000 rpm/IA20 rotor/10 min] and dialyzed against buffer HEMG until the conductivity was equivalent to a reading obtained with HEMG, 0.12 M KCl (approximately 3 hr). Precipitates were removed by centrifugation.
Fractionation of the embryo nuclear extract

Nuclear extract was applied to a heparin-Sepharose CL-6B column (Pharmacia, 1.3 ml extract/ml bed volume), equilibrated with HEMG, 0.1 mM KCl and washed with 3 column volumes of the same buffer. Protein was eluted with 2 column volumes of HEMG, 0.4 mM KCl, and the pooled fractions were dialyzed against HEMG, 40 mM KCl to a conductivity equivalent to that obtained with HEMG, 0.12 mM KCl. The dialyzed protein fraction (H.4, 8 mg/ml protein) was applied to a column of DEAE-Sepharose CL6B (Pharmacia, 1.5 ml H.4/ml bed volume), and the column was washed with 3 volumes of HEMG, 0.1 mM KCl, and eluted with 2 volumes of HEMG, 0.25 M KCl. Proteins flowing through the DEAE-Sepharose [D.1] or eluted with 0.25 M KCl [D.25] were pooled, aliquoted, frozen in liquid nitrogen, and stored at −80°C.

In vitro transcription

Run-off assays were performed essentially as described in Parker and Topol (1984a). In a final volume of 25 μl, 25 mM HEPES (pH 7.6), 6 mM MgCl₂, 50 mM KCl, 1 mM DTT, 0.05 mM EDTA, 3% glycerol, 500 μM each of ATP, CTP, and UTP, 50 μM GTP, 6 μCi of 410 Ci/mole [32P]GTP (Amersham), 4–20 μg/ml of linearized DNA template, and 40–160 μg crude nuclear extract were incubated for 30 min at 25°C. Reactions were stopped by addition of 4 μl of a stop mix consisting of 200 mM HEPES (pH 7.6), 10 mM EDTA, and unlabeled ATP, GTP, and UTP. After addition of 2 μl of an end-labeled, synthetic 20-nucleotide-long primer at a concentration of 0.5 μM and asp. act. of 30000 cpm/pmole primer. Two

DNA 1 protection assays

Footprint reactions were performed in a volume of 20 μl, containing 25 mM HEPES [pH 7.6], 50 mM KCl, 6 mM MgCl₂, 0.1 mM EDTA, 10–20 ng of end-labeled DNA fragment, and up to 10 μl of the various protein fractions. Reactions were assembled on ice in the absence of the DNA probe and allowed to equilibrate for 10 min. DNA probe was added and the reactions were incubated on ice for an additional 15 min. The reactions were then placed at room temperature, diluted twofold with a solution of 10 mM CaCl₂ and 20 mM MgCl₂ and digested with an appropriate amount of DNase I (DPRF, Worthington, 1.6 to 20 ng) for 1 min. The reaction was stopped with 100 μl stop mix (0.1% SDS, 200 μg/ml RNA carrier, 20 mM EDTA, 0.2 mM KCl). Reaction products were extracted with phenol/chloroform, precipitated with ethanol, and electrophoresed on 6% or 10% acrylamide (20 : 1 acrylamide/bisacrylamide)/7 M urea sequencing gels.

Cloned segments of Drosophila DNA used for in vitro transcription

DNA fragments used as templates for in vitro transcription included: a plasmid containing the Drosophila actin 5C gene in an 8.7-kb EcoRI–SalI segment of the plasmid pDmA2 (Fyrberg et al. 1980); the distal and proximal promoters for alcohol dehydrogenase (Adh) in a 4.7-kb EcoRI fragment from plasmid p13E-3 cloned into pUC13 (Heberlein et al. 1985); and the promoter for fushi tarazu in a 3.5-kb HindIII fragment.

Template for transcription of the engrailed gene were derived from plasmid p6:4.7, a 4.7-kb EcoRI fragment of phage E7 (Kuner et al. 1985). The template Δ − 887 to + 231 of the 4.7-kb EcoRI fragment cloned into pEMBL 18. Upstream deletion templates Δ − 130 and Δ − 118 were generated by the DNase I/MnCl₂ method (Hong 1982); they retain 130 and 118 bp, respectively, of upstream sequence and 231 bp of the engramed transcribed region. The remaining upstream deletion mutants were derived from Δ − 118, which was linearized at the upstream end with HindIII, digested with exonuclease III and nuclease S1, treated with T4 DNA polymerase to fill in the ends, and recircularized with T4 DNA ligase (Henikoff 1984). With the exception of Δ + 4 and Δ + 15, each mutant retains various lengths of upstream regions and 231 bp of the downstream transcribed region. Δ + 4 and Δ + 15 retain no upstream sequences and only 228 and 216 bp of transcribed region, respectively.

For a control template, a 12-bp BamHI linker was inserted at a filled DdeI site (position + 127) in Δ − 118. This control template, Δ − 118Bam, was also used to generate deletions from the downstream end. To introduce deletions, Δ − 118Bam was linearized with BamHI, treated with exonuclease III and nuclease S1, and treated with T4 DNA polymerase to fill in the ends. These fragments were cleaved with PstI at position + 231 and ligated to a purified BamHI/PstI fragment (engramed sequences + 127 to + 231) of Δ − 118Bam, followed by end filling and recircularization. This scheme reintroduced the transcribed region from + 127 to + 231, allowing use of primer 2 in the primer extension assays.

The internal deletion mutants Δ − 52/−34, Δ − 65/−34, Δ − 80/−38, and Δ − 77/−26 were derived from Δ − 118. This plasmid was linearized with HpaI at position − 38 and was digested with exonucleaseIII and nuclease S1 prior to end filling and recircularization. This procedure resulted in the removal of between 16 and 49 bp within a region bounded by positions − 80 and − 26.

To construct the engramed–CAT fusion templates, various lengths of the engramed promoter were joined to the polylinker.
directly upstream of the CAT coding sequences of plasmid pDM26 [Mismer and Rubin 1987] as follows: Δ−118DM26, with a BamHI fragment of Δ−118Bam containing 118 bp of upstream and 127 bp of downstream engrailed sequences into the BamHI site; Δ−960DM26, with a ClaI/BamHI-engrailed fragment (−960 to +127) into SalI/BamHI-cut pDM26; Δ−305DM26, Δ−38DM26, and Δ+2DM26 from Δ−960DM26 after digestion with Avai, HapI, and HincII nucleases, respectively. Δ−118DM26 derivatives with 39, 37, or 30 bp of downstream sequences were generated with XbaI–BamHI fragments from deletion mutants Δ+39, Δ+37, and Δ+30, respectively, and XbaI/BamHI-cut pDM26.

Transformations were carried out as described by DiNocera and Dawid [1983] using 10 μg of template DNA and 1 μg of the β-galactosidase control DNA per 60-mm tissue culture plate. Cells were harvested after 48 hr and assayed for CAT [Gorman et al. 1982] and β-galactosidase [Miller 1972] activity.

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