NELF and DSIF cause promoter proximal pausing on the hsp70 promoter in Drosophila

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NELF and DSIF collaborate to inhibit elongation by RNA polymerase IIa in extracts from human cells. A multifaceted approach was taken to investigate the potential role of these factors in promoter proximal pausing on the hsp70 gene in Drosophila. Immunodepletion of DSIF from a Drosophila nuclear extract reduced the level of polymerase that paused in the promoter proximal region of hsp70. Depletion of one NELF subunit in salivary glands using RNA interference also reduced the level of paused polymerase. In vivo protein–DNA cross-linking showed that NELF and DSIF associate with the promoter region before heat shock. Immunofluorescence analysis of polytene chromosomes corroborated the cross-linking result and showed that NELF, DSIF, and RNA polymerase IIa colocalize at the hsp70 genes, small heat shock genes, and many other chromosomal locations. Finally, following heat shock induction, DSIF and polymerase but not NELF were strongly recruited to chromosomal puffs harboring the hsp70 genes. We propose that NELF and DSIF cause polymerase to pause in the promoter proximal region of hsp70. The transcriptional activator, HSF, might cause NELF to dissociate from the elongation complex. DSIF continues to associate with the elongation complex and could serve a positive role in elongation.

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Inhibition of transcriptional elongation is an important step in regulating the expression of numerous genes in eukaryotic cells [Kim et al. 2001]. In the case of the hsp70 heat shock gene in Drosophila, RNA polymerase II [Pol II] is stably paused in the promoter proximal region before heat shock induction [Lis 1998]. Evidence of promoter proximal pausing has been obtained for a variety of other genes including the c-myc and c-fos genes [Krumm et al. 1992; Plet et al. 1995; Pinaud and Mirkovic 1998]. Law et al. [1998] provided evidence that the RNA polymerase associated with as many as 20% of genes in cultured Drosophila cells might be in a paused state.

Phosphorylation of Pol II appears to be a key step in converting Pol II from an elongation incompetent to elongation competent state [Weeks et al. 1993; O’Brien et al. 1994]. During transcription, extensive phosphorylation occurs at the C terminus of the largest subunit—a region called the CTD [Komarnitsky et al. 2000; Cho et al. 2001]. The CTD is composed of a 7-amino-acid sequence that is repeated 26 times in yeast, 42 times in Drosophila, and 52 times in human. Pol II associates with the promoter as an unphosphorylated form called PolIIa [Luet al. 1991]. In vitro, the CTD becomes highly phosphorylated during or shortly after transcriptional initiation [Laybourn and Dahmus 1990]. This hyperphosphorylated form of RNA polymerase is called PolIIo.

In vivo, the Pol II found paused on the hsp70 gene before heat shock induction is in the IIa state [Weeks et al. 1993; O’Brien et al. 1994]. Following heat shock, the polymerase detected on the body of the gene is in the hyperphosphorylated Pol IIo state. This suggests that the release of the paused polymerase is coupled to a hyperphosphorylation of the CTD. Consistent with this hypothesis is the finding that expression of a protein consisting of the Gal4 DNA-binding domain fused to the kinase P-TEFb activates transcription of an hsp70 promoter containing binding sites for the Gal4 protein [Lis et al. 2000]. Normally, hsp70 is induced during heat
shock by HSF. Although no direct association between P-TEFb and HSF is evident, P-TEFb associates with the hsp70 gene during heat shock [Lis et al. 2000]. There is mounting evidence that several transcriptional activators function in part by mediating phosphorylation of the CTD either through the kinase P-TEFb or the general transcription factor TFIIH [Blau et al. 1996; Barboric et al. 2001; Eberhardy and Farnham 2001].

An understanding of how phosphorylation of the CTD may control transcriptional elongation has emerged from the biochemical analysis of the sensitivity of transcription toward the nucleoside analog DRB [Yamaguchi et al. 1998]. DRB inhibits kinases that are responsible for phosphorylating the CTD, and it causes greater than 50% inhibition of Pol II transcription in human cells [Sehgal et al. 1976; Mancebo et al. 1997; Chao and Price 2001]. DRB appears to inhibit elongation as cells are seen to accumulate short transcripts and suppress production of long transcripts [Tamm and Kikuchi 1979; Chodosh et al. 1989]. Elongation by purified Pol II is not inhibited by DRB, indicating that DRB targets other proteins that modulate elongation [Kephart et al. 1992]. The kinase P-TEFb was first identified as a factor required to reconstitute DRB-sensitive transcription [Marshall and Price 1995]. More recently, two other proteins called DSIF and NELF were discovered to be required for DRB inhibition [Wada et al. 1998a; Yamaguchi et al. 1999a]. The results of immunodepletion experiments support a model in which DSIF and NELF inhibit elongation by Pol IIa, and P-TEFb relieves this inhibition by phosphorylating the polymerase [Wada et al. 1998b; Yamaguchi et al. 1999a]. Additional support for this model comes from the observation that NELF and DSIF only inhibit elongation by Pol IIa and not by Pol IIo in a well-defined elongation reaction [Yamaguchi et al. 1999a].

Chromatin immunoprecipitation analysis has revealed that DSIF is associated with the hsp70 promoter before heat shock induction, so it is reasonable to postulate that this factor is involved in promoter proximal pausing [Andrulis et al. 2000]. There is also strong evidence that DSIF stimulates transcriptional elongation, however. Immunofluorescence analysis of polytene chromosomes in Drosophila shows that DSIF associates with strongly transcribed regions of the genome [Andrulis et al. 2000; Kaplan et al. 2000]. DSIF stimulates tat-dependent elongation in reconstituted transcription reactions apparently by counteracting premature termination and transcriptional arrest [Wu-Baer et al. 1998; Bourgeois et al. 2002]. DSIF is also known as Spt4 and Spt5, which correspond to its small and large subunits. Several alleles of spt5 exhibit synthetic lethality when combined with mutations in Pol II or TFIIIS, a stimulatory elongation factor [Hartzog et al. 1998]. At present, there is no clear explanation for how DSIF functions both as a positive and a negative elongation factor. We have investigated the potential roles of DSIF and NELF in regulating promoter proximal pausing at the hsp70 promoter in Drosophila. We provide both physical and functional data supporting the hypothesis that NELF and DSIF are important for pausing Pol II in the promoter proximal region. Moreover, the mechanism of transcriptional activation appears to involve ejection of NELF from the paused Pol II.

Results

**Immunodepletion of DSIF from a Drosophila nuclear extract inhibits promoter proximal pausing in vitro**

Previously, we showed that promoter proximal pausing could be reconstituted in Drosophila nuclear extracts [Li et al. 1996; Benjamin and Gilmour 1998]. The paused polymerase was detected by monitoring the reactivity of the transcribed DNA toward permanganate, which preferentially oxidizes thymines in single-stranded regions. When nuclear extract was incubated with the hsp70 promoter in the absence of nucleotides, very little permanganate reactivity was detected in the region 20–40 nucleotides downstream from the transcription start (Fig. 1A, lane 1). When all four nucleotides were present, the permanganate reactivity of the thymines at +22 and +30 was enhanced for a mock-depleted extract, which still contains DSIF (Fig. 1A, lane 3). Immunodepletion of DSIF from the nuclear extract decreased the level of permanganate reactivity detected at +22 and +30 (Fig. 1A, cf. lanes 5 and 3). This decrease could be partially reversed by supplementing the depleted extract with a protein fraction that had been enriched 25-fold for Drosophila DSIF (Fig. 1A, lane 7). As shown by the immunoblot in Figure 1B, the amount of partially purified DSIF was sufficient to return the level of DSIF to normal (cf. lanes 1 and 3).

The decrease in permanganate reactivity resulting from immunodepletion of DSIF could be caused by either a decrease in pausing or a decrease in initiation. To determine if the depletion of DSIF affected initiation, GTP in the reaction was replaced with 3′ O-methyl GTP. 3′ O-methyl GTP halts elongation when Pol II reaches +15, where the first guanine nucleotide is incorporated into the nascent transcript. The level of permanganate reactivity at +22 was essentially the same for the mock-depleted, DSIF-depleted, and the rescued extracts (Fig. 1A, lanes 2,4,6). Previous analysis demonstrated that our reaction conditions allow only one round of initiation [Benjamin and Gilmour 1998]. Therefore, the level of initiation on the hsp70 promoter in the three situations is similar. The decrease in permanganate reactivity at +22 and +30 caused by the depletion of DSIF is therefore attributable to inhibition of promoter proximal pausing.

**Identification of NELF-D and NELF-E from Drosophila**

We were motivated to investigate Drosophila NELF because biochemical analyses indicated that both DSIF and NELF were required to repress elongation in extracts from human cells [Yamaguchi et al. 1999a]. BLAST analyses identified single candidates for NELF-D and NELF-E in Drosophila (data not shown). These genes have been designated CG9984 and CG5994 by the Berke-
LEY Drosophila Genome Project. Drosophila NELF-D has 54% identity over its entire length with human NELF-D. 40% of the amino acids from 1 to 107 and 37% of the amino acids from 140 to 277 of Drosophila NELF-E are identical to human NELF-E. Amino acids 165–236 in Drosophila NELF-E are homologous to the RNA-binding domain of human NELF-E. This region is essential for NELF to inhibit elongation in vitro (Yamaguchi et al. 2002).

The N-terminal portion of Drosophila NELF-D (amino acids 33–154) and the entirety of NELF-E were expressed and purified from Escherichia coli, and each polypeptide was used to raise antibody. A Western blot analysis showed that each antiserum but not the corresponding preimmune serum recognized a polypeptide of the expected size in a Drosophila nuclear extract [Fig. 2A]. As further evidence that Drosophila NELF-D and NELF-E

Figure 1. Immunodepletion of DSIF from a Drosophila nuclear extract inhibits promoter proximal pausing. [A] In vitro transcription reactions were performed with extracts that had been subjected to depletions with antibody against DSIF or against an irrelevant protein (Mock depleted). Paused polymerase was detected by permanganate footprinting. Lane 1 shows the results for a transcription reaction performed in the absence of nucleotides. Lanes 3, 5, and 7 were performed in the presence of all four nucleotides. Lanes 2, 4, and 6 were performed with 125 µM 3′ O-methyl GTP substituted for GTP. [B] Immunoblot analysis for DSIF in extracts used in the reconstituted transcription reactions. Lanes 1 and 2 are 2-µL portions of mock-depleted or DSIF-depleted extract. Lane 3 is a mixture of 2 µL of DSIF-depleted extract and 2 µL of the partially purified DSIF, which represents the proportions of proteins used in the transcription reaction used for lanes 6 and 7 in A.

Figure 2. Identification of Drosophila NELF-D and NELF-E. [A] Immunoblot analysis shows that antibodies raised against the Drosophila homologs bind specifically to polypeptides of the sizes predicted for the respective cDNA clones. Nuclear extract was run on SDS-PAGE and blotted to nitrocellulose. Individual nitrocellulose strips harboring nuclear proteins were incubated with preimmune serum for NELF-D or NELF-E [lanes 1, 3, respectively] or with antiserum for NELF-D or NELF-E [lanes 2, 4, respectively]. The NELF-D antiserum specifically binds a polypeptide of ∼60 kD and the NELF-E antiserum specifically binds a polypeptide of ∼36 kD. [B] Coimmunoprecipitation data indicates that Drosophila NELF-D and NELF-E associate with each other. Nuclear extract containing 150 mM KCl was incubated with preimmune antiserum [lanes 1, 2], affinity-purified NELF-D antibody [lanes 3,4], NELF-D antiserum [lanes 5,6], NELF-E preimmune antiserum [lanes 7,8], or NELF-E antiserum [lanes 9,10]. Immune complexes were recovered with protein A sepharose, and the presence of NELF-D and NELF-E was assessed by immunoblot analysis. S, supernatant; P, immunoprecipitated material.
are homologs to human NELF subunits, NELF-D and NELF-E were found to associate with each other in *Drosophila* nuclear extract. Antibodies against either polypeptide coprecipitated the other polypeptide. Affinity-purified anti-NELF-D antibody or crude antiserum against NELF-D precipitated both NELF-D and NELF-E [Fig. 2B, lanes 4,6]. Crude antiserum against NELF-E also precipitated both NELF-D and NELF-E [Fig. 2B, lane 10]. In contrast, preimmune serum for either NELF-D or NELF-E failed to precipitate either protein [Fig. 2B, lanes 2,8].

**NELF-D, NELF-E, DSIF, and GAGA factor associate with the hsp70 promoter region before heat shock induction**

Previously, Andrulis et al. (2000) used chromatin immunoprecipitation to determine that DSIF associates with the *hsp70* promoter before heat shock induction. The results in Figure 3 show that NELF-D and NELF-E also associate with the *hsp70* promoter. *Drosophila* cells were treated with formaldehyde to form protein–protein and protein–DNA cross-links, lysed in SDS, sonicated to shear DNA to sizes from 200 to 700 bp, and subjected to immunoprecipitations. DNA recovered from each immunoprecipitate was then analyzed for the presence of DNA corresponding to the promoter region [5′] or the 3′ region of the *hsp70* gene (Fig. 3A).

Figure 3B provides representative results from an analysis, and the graph quantifies the amounts of DNA recovered from the 5′ and 3′ ends of the *hsp70* gene that were recovered from various immunoprecipitates for two independent experiments. The amount of material recovered from all of the preimmune precipitates averaged 0.031% with a standard deviation of 0.015 and a maximum value of 0.068%. This range establishes the background inherent to the technique. In cells, the level of NELF-D, NELF-E, DSIF, and GAGA factor cross-linking to the 5′ region significantly exceeded the level cross-linking to the 3′ region of the gene. Approximately 0.4% of the 5′ region cross-linked to NELF-D and NELF-E, and 2.5% of the 5′ region cross-linked to DSIF and GAGA factor. In contrast, the amount of DNA from the 3′ region cross-linking to any of these proteins was in the range of background. Therefore, all four of these proteins appear to preferentially associate with the promoter region before heat shock induction. Previous UV cross-linking and genomic footprinting analyses showed that GAGA factor associates with the promoter region before heat shock (O’Brien et al. 1995; Weber et al. 1997). The detection of DSIF on the promoter region confirms results reported by Andrulis et al. (2000). The results in Figure 3 show for the first time that NELF also associates with the promoter region.

**Depletion of NELF-E in salivary glands reduces the level of paused polymerase found on hsp70 in live cells**

We tested the hypothesis that NELF is involved in promoter proximal pausing by using RNA interference to deplete NELF-E from salivary glands. Interference RNA against NELF-E was shown recently to cause developmental defects or lethality depending on where the interference RNA was expressed (Enerly et al. 2002). For our experiment, one transgenic fly line, 17A10, contained a transgene encoding an interference RNA against NELF-E that was controlled by Gal4p. A second fly line, 1824, contained a transgene that expressed Gal4p in salli-

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**Figure 3.** Chromatin immunoprecipitation analysis of NELF, DSIF, and GAGA factor. *Drosophila* Schneider 2 cells were subjected to formaldehyde cross-linking. Sonicated detergent lysates from the cells were immunoprecipitated with preimmune serum [Pre-D or Pre-E] or with antiserum against NELF-E, NELF-D, DSIF, or GAGA factor. DNA was purified from each immunoprecipitate, and the presence of DNA from the 5′ and 3′ portions of the *hsp70* gene was detected by PCR. Various amounts of material representing the input were amplified to quantify the amount of DNA recovered from the immunoprecipitates. [A] A schematic of the *hsp70* transcription unit and the regions that were amplified. [B] The results from one of two independent experiments that were done. The bars represent radiolabeled PCR products that were detected after the DNA was run on an acrylamide gel and the gel was subjected to analysis with a PhosphorImager. Because preliminary experiments showed that DSIF and GAGA factor cross-linked more efficiently than NELF-D and NELF-E, 5× less of the DNA recovered from the DSIF and GAGA factor immunoprecipitates were used in the PCR reactions. The graph shows a compilation of the results from two independent experiments. The height of each bar shows the average percent of input material detected in immunoprecipitates from two experiments, and each error bar shows the range of measurements.
vary glands (Brand et al. 1994). The two lines were mated, and salivary glands derived from the progeny and the parental lines were subjected to molecular analysis. Immunoblot analysis of whole salivary gland lysates revealed that RNAi against NELF-E significantly diminished the amount of NELF-E in the glands (Fig. 4A). Interestingly, the level of NELF-D was unaffected. Genomic footprinting with permanganate was performed to determine if depleting NELF-E caused a decrease in promoter proximal pausing on hsp70. Previously, we showed that the paused Pol II causes thymines at +22 and +30 on the nontranscribed strand to be hyperreactive to permanganate oxidation in salivary glands (Weber et al. 1997). Salivary glands were dissected from third-instar larvae and treated with permanganate. DNA was isolated, and the pattern of permanganate reactivity was analyzed following cleavage at oxidized thymines and ligation-mediated PCR. As shown in Figure 4B, the permanganate reactivity at +22 and +30 was less in the RNAi glands than in glands from either parental line. This affect was observed for three independent RNAi lines [data not shown].

Immunofluorescence analysis of NELF-D on polytene chromosomes from Drosophila salivary glands

Immunofluorescence analysis of polytene chromosomes was employed to corroborate the results of chromatin immunoprecipitation and to learn more about the genome-wide distribution of NELF. We focused on NELF-D because we were able to affinity-purify NELF-D antibody that stained the polytene chromosome. Affinity-purified antibody against NELF-D broadly stains polytene chromosomes [Fig. 5A]. No staining was observed with preimmune serum or with material isolated in a mock affinity purification of antibody [data not shown]. Hoechst dye stains DNA and stains polytene chromosomes in a pattern of bands that coincides closely with the pattern of bands on chromosomes typically detected by phase contrast microscopy [Fig. 5B]. NELF-D was restricted to the regions known as interbands, which lie between the bands of DNA staining [Fig. 5C,D].

Several regions of the polytene chromosomes consistently stained more intensely than other regions of the chromosomes. These regions include 63A, 67B, 87C, and 89B, which are also evident for independent samples shown in Figure 6. The staining at 87C was of interest because at least three copies of the hsp70 gene reside at this locus. Staining at 67B is also noteworthy because four small heat shock genes are located in this region, and three have been shown to harbor paused polymerase (Lis 1998).

A band of fluorescence on a polytene chromosome could encompass many genes and several hundred kilobases of DNA. To test if the fluorescence at 87C could be caused by association between NELF-D and the hsp70 promoter, we compared chromosomes from larvae that had a copy of the hsp70 promoter inserted at 87E to chromosomes lacking this insertion. The hsp70 promoter had been inserted as part of a P-element construct, and it consisted of nucleotides −194 to +84 (Wu et al. 2001). Genomic footprinting with permanganate showed that a paused polymerase was present on this promoter [data not shown]. Inspection of 20 different chromosomes derived from three different slides for each type of larvae clearly showed that the staining for NELF-D at 87E was enhanced by the presence of the hsp70 promoter. Repre-

Figure 4. NELF-E RNA interference inhibits synthesis of NELF-E in salivary glands and inhibits pausing on the hsp70 gene. Fly line 1824 expresses Gal4 in salivary glands. Fly line 17A10 contains a Gal4-activated NELF-E RNAi construct. RNAi is the designation for progeny generated by mating 1824 and 17A10. [A] Immunoblot analysis of NELF-D and NELF-E in whole salivary gland lysates. [B] Permanganate genomic footprinting analysis of hsp70 in salivary glands isolated from non-heat-shocked larvae. Lanes 1 and 5 are derived from genomic DNA partially cleaved at purines. (Lane 2) DNA derived from 1824 salivary glands that were treated with permanganate. (Lane 3) DNA derived from RNAi salivary glands that were treated with permanganate. (Lane 4) DNA derived from 17A10 glands that were treated with permanganate.
sentative examples are shown in Figure 5E. The band marked with a white dot serves as a reference. NELF-D staining exceeds the intensity of staining at the reference only when the transgenic promoter is present. In addition, NELF-D staining of 87A is evident in all samples where the chromosomes are well spread in this region. Two copies of hsp70 reside at 87A. Taken together, these data corroborate the results of the ChiP analysis showing that NELF associates with the hsp70 promoter before heat shock induction.

Comparison of the distributions of NELF-D and DSIF

We compared the distributions of NELF-D and DSIF because NELF and DSIF collaborate to inhibit transcriptional elongation in vitro (Yamaguchi et al. 1999a). Double-label immunofluorescence reveals that both proteins are broadly distributed among interband regions [Fig. 6A,B]. The intensity of staining for DSIF, however, is much more varied than the staining for NELF-D. The sites staining very strongly for DSIF (3C, 10E, 90B, 68C, and 71E) are all sites registered as puffs in third-instar larvae and are regions of high transcriptional activity (Ashburner 1972). Notably, NELF-D does not colocalize with DSIF in these highly transcribed regions.

Outside the regions of intense DSIF staining, there is significant colocalization of NELF-D and DSIF. The relationship between the intensities of fluorescence and the quantities of protein are not known. Therefore, based solely on the intensity of fluorescence, we cannot be certain which regions of the chromosome contain similar amounts of the two proteins. Nevertheless, we observe that the relative intensity of staining for both NELF-D and DSIF at 87A, 87C, and 87E are similar. Moreover, when the intensities of the two colors were adjusted so the merged image appeared yellow at these three loci, 67B also appeared yellow (Fig. 6C). As these loci all harbor paused Pol II, these results are consistent with the hypothesis that these locations contain amounts of NELF-D and DSIF that cause pausing.

Figure 5. Immunolocalization of NELF-D on polytene chromosomes from larvae. Polytene chromosome spreads were prepared from salivary glands of third-instar larva. [A] The chromosomes were incubated with affinity-purified NELF-D antibody and then incubated with Alexa 568 conjugated secondary antibody. [B] DNA was detected by staining with Hoechst. [C] The merged image shows the NELF-staining pattern in red and the DNA staining pattern in blue. [D] A split image of the end of chromosome 2L with the DNA stain on the top half and NELF-D staining on the bottom half. [E] Representative examples of the NELF-D staining pattern in the region encompassing the endogenous hsp70 genes and an hsp70 transgenic promoter that is inserted at 87E. The dot marks a band that serves as a reference for comparing the intensity of staining at 87E where the transgenic promoter resides. The top three pictures are from chromosomes containing the insert and the bottom three pictures are from chromosomes that lack the insert. Note that the intensity of staining for NELF-D is stronger than the reference in cases containing the transgenic promoter, but not for the cases lacking the transgenic promoter.
Figure 6. Immunolocalization of NELF-D, DSIF, and Pol IIa. Chromosome spreads from larvae containing an *hsp70* transgenic promoter at 87E were stained with either a combination of affinity-purified rabbit anti-NELF-D antibody and guinea pig anti-DSIF antibody (A–C) or a combination of affinity-purified rabbit anti-NELF-D antibody and a monoclonal antibody (8WG16) against the unphosphorylated CTD of Pol IIa (D–F). The chromosomes were also stained with Hoechst to assist in mapping locations on the chromosomes (data not shown). (A) The NELF-D antibody visualized with Alexa 568 conjugated anti-rabbit antibody. (B) The DSIF antibody visualized with Alexa 488 conjugated anti-guinea pig antibody. (C) A merge with red representing NELF-D and green representing DSIF. (D) The NELF-D antibody visualized with Alexa 568 conjugated anti-rabbit antibody. (E) The Pol IIa monoclonal antibody visualized with Alexa 488 conjugated anti-mouse antibody. (F) A merge with red representing NELF-D and green representing Pol IIa.
Comparison of the distributions of NELF-D and Pol IIa

Pol II paused on the hsp70 promoter before heat shock is in the Pol IIa state (Weeks et al. 1993; O’Brien et al. 1994). The results in Figure 6D and E show that NELF-D and Pol IIa localize to interbands, but as was observed for DSIF, the intensity of staining for Pol IIa is much more varied than the staining for NELF-D. The regions most strongly stained for Pol IIa correspond to highly transcribed regions (71E, 68C, 90B, 3C). Although these regions are known to be enriched in Pol IIo, Weeks et al. (1993) reported previously that antibody against Pol IIa strongly stained highly transcribed regions. Some heptamers constituting the CTD may remain unphosphorylated, or the antibody may weakly bind heptamers phosphorylated on Ser 5 (Cho et al. 2001). Close examination of 87E revealed that Pol IIa staining was enhanced in cases when the hsp70 transgene was present but not in those cases when it was absent (data not shown).

DSIF but not NELF accumulates on the hsp70 gene following heat shock induction

We were intrigued to find that antibody against NELF-D failed to stain the highly transcribed regions on polytene chromosomes that were marked by strong staining against DSIF or Pol IIa (Fig. 6C,F). Perhaps NELF-D dissociates from the Pol II when an activator causes release of paused Pol II. To test this hypothesis, we determined if NELF-D associates with the hsp70 genes during heat shock induction. A 20-min heat shock induces transcription of the heat shock genes and causes 87A and 87C to become puffed (Fig. 7). 87E also becomes puffed in the case where an hsp70 transgene is inserted. The antibodies against DSIF or Pol IIa strongly stained 87A and 87C in accord with previous reports (Andrulis et al. 2000; Kaplan et al. 2000), and 87E when the hsp70 transgenic promoter was present (Fig. 7A,C). In contrast, NELF-D antibody only weakly stained puffed regions. Therefore, NELF-D appears to dissociate from Pol II during active transcriptional elongation.

Discussion

Many proteins have been identified that stimulate transcriptional elongation possibly by alleviating barriers to elongation (Shilatifard 1998). TFIIF, TFIIS, and elongin can stimulate elongation by purified polymerase and function in part to overcome intrinsic features of the nucleic acid sequence that seem to interfere with the movement of Pol II. FACT, elongator, and Swi/Snf seem to function to alleviate barriers to elongation imposed by the chromatin structure. Whether or not these factors are used in gene-specific ways to control gene expression is not known. Evidence for activator-dependent control of elongation has only been provided for Swi/Snf (Brown et al. 1996; Sullivan et al. 2001). Many of these elongation factors could represent part of the repertoire of elongation factors that is called into action at any gene depending on what barriers the polymerase encounters. In contrast, our results provide evidence that DSIF and NELF could be at the heart of a widely used mechanism for implementing gene-specific control of expression.

We have taken a multifaceted approach to provide both physical and functional evidence that NELF and DSIF participate in promoter proximal pausing. Chromatin immunoprecipitation analysis provided physical evidence that NELF and DSIF are associated with the hsp70 promoter region before heat shock. This was corroborated by immunofluorescence analysis of polytene chromosomes. Two types of functional data indicate that NELF and DSIF participate in promoter proximal pausing on hsp70. Immunodepletion of DSIF caused a decrease in the level of paused polymerase that was reconstituted on the hsp70 promoter in nuclear extracts. At-
temps to determine if immunodepleting NELF from the nuclear extract affects pausing in vitro have been marginally successful (data not shown). We observed a slight decrease in the permanganate reactivity at $+22$ after depleting NELF, but Western blot analysis indicated that the NELF depletions were incomplete. We have successfully implemented an RNAi approach to evaluate the contribution of NELF toward pausing on hsp70 in live cells. The permanganate reactivity of thymines in the promoter proximal region of the RNAi salivary glands is twofold less than the reactivity observed for either parent. Note that genomic footprinting with permanganate is a very sensitive way to detect relative levels of polymerase associated with the DNA in salivary glands [Tang et al. 2000]. Therefore the decrease in permanganate reactivity caused by the RNAi indicates that there is less paused polymerase. The loss of pausing in RNAi flies was incomplete. A low level of NELF-E persists in spite of the interfering RNA [data not shown]. The residual NELF-E may be sufficient for pausing to occur because the paused state appears to be very stable once formed [see below].

Figure 8 provides a model for the control of promoter proximal pausing on the hsp70 promoter. DSIF and NELF have a central role in the pausing process by associating with Pol IIa during elongation. The generation of the paused polymerase is preceded by steps that participate in chromatin remodeling and initiation. GAGA factor associates with the hsp70 promoter before heat shock induction [Giardina et al. 1992; O’Brien et al. 1995; Wei-

ber et al. 1997]. GAGA factor interacts with the chromatin remodeling factor, NURF, and could target chromatin remodeling activity to the promoter [Tsukiyama et al. 1994; Xiao et al. 2001; Badenhorst et al. 2002]. A pre-initiation complex would then assemble. It is not clear what proteins in addition to components of the general transcription machinery participate in assembly of the initiation complex.

We propose that promoter proximal pausing occurs when the nascent transcript emerges from the RNA exit channel of the Pol II and is grabbed by the NELF-E subunit. Tethering of the NELF-E to the elongation complex would generate a rigid body that could restrict the movement of the Pol IIa. This model is supported by several observations. The paused polymerase is in the Pol IIa state [O’Brien et al. 1994], and NELF and DSIF only inhibit elongation by Pol IIa [Yamaguchi et al. 1999a]. In vitro transcription analysis indicates that the elongation complex is not receptive to inhibition by NELF and DSIF until the nascent transcript is ~30 nucleotides long [Yamaguchi et al. 1999a]. This length coincides approximately to the distance polymerase elongates on hsp70 before it pauses [Rasmussen and Lis 1993]. In vitro transcription analyses indicate that DSIF and NELF associate with polymerase shortly after initiation but probably before the polymerase reaches the region of pausing [Ping and Rana 2001; Bourgeois et al. 2002]. Finally, NELF-E has an RNA-binding motif that is essential for its inhibitory action [Yamaguchi et al. 2002].

Although NELF and DSIF are sufficient to slow the elongation rate of purified Pol IIa, we suspect additional proteins are involved in stably pausing Pol II on the hsp70 promoter. In cell-free transcription reactions done with other promoters, the pausing caused by DSIF and NELF appears to be transient—the polymerase eventually moves forward if given enough time [Wada et al. 1998a]. In contrast, several observations indicate that the Pol II on hsp70 is stably paused. The paused Pol II remains associated with the hsp70 promoter when nuclei are isolated from uninduced cells, and sarkosyl or high salt must accompany addition of nucleotides to cause the Pol II to resume elongation [Rougvi and Lis 1988]. In a cell-free system, Pol II remains stably paused on the hsp70 promoter for at least 30 min [Li et al. 1996]. GAGA factor might be involved in stabilizing the pause because mutations in the GAGA element result in a loss of paused Pol II [Lee et al. 1992].

Heat shock rapidly induces transcription as a result of the association of HSF with sites located upstream from the TATA element [Jedlicka et al. 1997]. Our data suggest that HSF may activate transcription in part by causing NELF to dissociate from the Pol II. How HSF might cause the release of NELF is unclear. Phosphorylation of Pol IIa is likely to be an important step as the Pol II found in the body of the gene during heat shock is hyperphosphorylated [Weeks et al. 1993]. Phosphorylation of DSIF is another possibility as this has been observed to occur early in elongation in vitro [Bourgeois et al. 2002]. It is also unclear which kinase might be responsible for phosphorylating the Pol II. P-TEFb is a candidate because it
associates with the hsp70 gene during heat shock induction, and HSF can be bypassed by directing a Gal4/P-TEFb fusion protein to the hsp70 promoter [Lis et al. 2000]. No interaction, however, has been detected between P-TEFb and HSF [Lis et al. 2000]. Recent results show that HSF associates with the mediator [Park et al. 2001b]. Drosophila mediator contains a kinase that phosphorylates the CTD, and phosphorylation can occur synergistically with the TFIIBH kinase [Park et al. 2001a]. Perhaps HSF recruits the mediator and in turn the mediator releases the paused polymerase by phosphorylating the CTD.

The strong immunofluorescence staining observed for DSIF at heat shock loci during heat shock indicates that DSIF is associated with many of the polymerase molecules transcribing the gene. RNA polymerase initiates at a rate of once every few seconds during heat shock resulting in a train of elongation complexes traversing the gene [Gilmour and Lis 1985]. In the absence of NELF, DSIF might act as a positive elongation factor. Shortly after DSIF was discovered, another investigation identified DSIF as a cofactor required for reconstituting Tat-dependent transcription [Wu-Baer et al. 1998]. In this situation, DSIF appears to be stimulating elongation. DSIF has been found in a complex with another positive elongation factor called Tat-SF1 [Kim et al. 1999; Parada and Roeder 1999]. Tat-SF1 was first identified as a stimulatory factor for Tat, but subsequent results indicate that Tat–SF1 may promote elongation on cellular genes [Li and Green 1998]. A hypothesis that unites the positive and negative activities of DSIF is to consider this factor an adaptor that connects other modulators to the elongation complex. In this regard, DSIF has been shown to bind on its own to Pol II [Yamaguchi et al. 1999b], whereas the stable association of NELF with Pol II requires the presence of DSIF [Yamaguchi et al. 2002]. NELF and DSIF appear to associate with several hundred interbands in polytene chromosomes. Each interband could contain many genes. The weak staining of interbands by Hoechst suggests that the DNA in the interbands is in a decondensed state. Residing in these decondensed regions could be genes whose primary control by band and genes whose primary control depends on circumstances [Hartzog et al. 1998]. A hypothesis that unites the positive and negative activities of DSIF is to consider this factor an adaptor that connects other modulators to the elongation complex. In this regard, DSIF has been shown to bind on its own to Pol II [Yamaguchi et al. 1999b], whereas the stable association of NELF with Pol II requires the presence of DSIF [Yamaguchi et al. 2002]. NELF and DSIF appear to associate with several hundred interbands in polytene chromosomes. Each interband could contain many genes. The weak staining of interbands by Hoechst suggests that the DNA in the interbands is in a decondensed state. Residing in these decondensed regions could be genes whose primary control mechanism does not involve a disruption of chromatin structure or even assembly of the initiation complex. Instead, alleviating repression by NELF and DSIF could underlie the mechanism of activation.

Materials and methods.

Immunodepletion analysis of nuclear extracts

Drosophila embryo nuclear extract was prepared as described by Biggin and Tjian [1988] except the final ammonium sulfate precipitate was dissolved in 60% the prescribed volume of buffer C modified to contain 0.45 M KCl. The extract was dialyzed against buffer C modified to contain 0.45 M KCl until the conductivity of the extract was equivalent to 0.5 M KCl/HEMG. HEMG is 25 mM HEPES at pH 7.6, 12.5 mM MgCl₂, 0.1 mM EDTA, 10% glycerol, 1 mM DTT, and 0.1 mM PMSF. The extract was clarified by centrifugation and stored at ~70°C before depletion. Guinea pig antiserum [200 µL] raised against the largest subunit of DSIF (dSF5) or an irrelevant protein were covalently coupled to 200 µL of protein A sepharose using dimethylpimelimidate [Harlow and Lane 1999]. For each depletion, 600 µL of nuclear extract was incubated at 4°C for ~3 h with 100 µL of antibody-sepharose. The extract was separated from the sepharose by a brief centrifugation and then incubated with a fresh 100 µL portion of antibody-sepharose. While the second cycle of depletion was occurring, the first portion of antibody-sepharose was washed several times with 0.5 M KCl/HEMG, followed by 0.1 M glycine at pH 2.5, followed by 0.5 M KCl/HEMG. Nuclear extract recovered from the second cycle of depletion was depleted a third time using the regenerated portion of antibody-sepharose. The extract from the third cycle of depletion was dialyzed for 2.5 h against 0.14 M KCl/HEMG. Protein concentrations were determined to be 19 mg/mL for both extracts. Immunoblot analysis indicated that at least 90% of the DSIF was removed from the extract.

Partial purification of DSIF

Nuclear extract from ~100 grams of embryos was prepared as described above except that the final ammonium sulfate pellet was dissolved in 0.7 M KCl/0.01% NP-40/HEMG and dialyzed against the same solution until the conductivity was equivalent to 0.75 M KCl/0.01% NP-40/HEMG. Then, 14.7 mL of extract was fractionated on a 1.8-L Sephacryl-300 column. Peak fractions of DSIF were identified by immunoblotting, pooled, and dialyzed against 0.1 M KCl/0.01% NP-40/HEMG. The pooled fraction was loaded onto an 8-ml MonoS column and eluted with a 160-mL gradient from 0.1 to 0.3 M KCl. Pooled fractions containing DSIF were loaded onto a 1-ml MonoQ column. The column was washed with 0.2 M KCl/0.01% NP40/HEMG, and DSIF was eluted with 0.3 M KCl/0.01% NP40/HEMG. The total yield of DSIF was ~4% and the DSIF was enriched ~25-fold.

Permanganate analysis of paused polymerase in reconstituted transcription reactions

Ten microliters of immunodepleted extract was combined with 10 µL of 0.15 M KCl/0.01% NP-40/HEMGN or 10 µL of partially purified DSIF and the mixture was maintained on ice. Transcription reactions were performed as described previously [Li et al. 1996; Benjamin and Gilmour 1998]. After a 30-min incubation at 22°C, samples were subjected to permanganate analysis as described previously [Benjamin and Gilmour 1998].

Cloning and production of antibodies

A BLAST search of the Drosophila genome using the sequence for human NELF-D identified EST clone LD42626. LD42626 DNA was obtained from the Berkeley Drosophila Genome Project, and sequencing showed that the clone contained an open reading frame encoding a 579 amino acid polypeptide with homology to human NELF-D. A primer set, 5’-ATATGCAGGAATGCCTGGAGAAGTTCC-3’ and 5’-GGCGATCCATATTTATCCTGAGGTG-3’, was used to amplify the region of LD42626 encoding amino acids 33–154. The amplified DNA fragment was digested with Ndel and BamHI, and then subcloned into the pET-28-[a] expression vector [Novagen]. The resulting 6x-His–tag fusion protein was expressed in BL21(DE3) cells and purified on a Ni-chelated column. Purified recombinant 6x-His–NELF-D protein was injected into rabbits to generate antibody.
Brief centrifugation and

Ten milliters of Chromatin immunoprecipitation analysis detected by immunoblotting. antibody beads were washed three times with 1-mL aliquots of KCl/HEMG, and the diluted extract was incubated with anti-

5×10^6/mL in T-flasks at 25°C in Schneider’s Drosophila medium (GIBCO-BRL) supplemented with 10% heat-inactivated fetal bovine serum. Formaldehyde cross-linking was done by adding 675 µL of 37% formaldehyde stock solution directly to the culture medium in the flask giving a final concentration of 1%. Cells were incubated at room temperature for 10 min, dislodged from the surface of the T-flask, transferred to centrifuge tubes on ice, and collected by centrifugation for 10 min. The antibody beads were washed three times with 1-mL aliquots of 0.15 M KCl/HEMG. The antibody beads were suspended in 100 µL of 1× SDS-PAGE buffer. Ten microliters of the supernatant and dissolved immunoprecipitate were run on 4%–15% polyacrylamide gradient SDS-PAGE, and NELF-D and NELF-E were detected by immunoblotting.

Immunoprecipitation of NELF from crude nuclear extracts

One hundred microliters of serum was incubated with 20 µL of 50% protein A sepharose beads at 4°C for 1 h. The beads were washed three times with 1 mL 0.15 M KCl/HEMG and most of the liquid was removed after each wash. Ten microliters of Drosophila embryo nuclear extract was mixed with 90 µL of 0.15 M KCl/HEMG, and the diluted extract was incubated with antibody beads for 2 h at 4°C. The antibody beads were collected by brief centrifugation and ~100 µL of supernatant was transferred to a fresh tube containing 100 µL of 2× SDS-PAGE buffer. The antibody beads were washed three times with 1-mL aliquots of 0.15 M KCl/HEMG. The antibody beads were suspended in 100 µL of 1× SDS-PAGE buffer. Ten microliters of the supernatant and dissolved immunoprecipitate were run on 4%–15% polyacrylamide gradient SDS-PAGE, and NELF-D and NELF-E were detected by immunoblotting.

Chromatin immunoprecipitation analysis

Ten milliters of Drosophila S2 cells were grown to a density of 5 × 10^6/mL in T-flasks at 25°C in Schneider’s Drosophila medium (GIBCO-BRL) supplemented with 10% heat-inactivated fetal bovine serum. Formaldehyde cross-linking was done by adding 675 µL of 37% formaldehyde stock solution directly to the culture medium in the flask giving a final concentration of 1%. Cells were incubated at room temperature for 10 min, dislodged from the surface of the T-flask, transferred to centrifuge tubes on ice, and collected by centrifugation at 700g for 10 min. Subsequent steps were done according to the chromatin immunoprecipitation procedure provided by Upstate Biotechnology. The immunoprecipitated DNA recovered from 5 × 10^6 cells was dissolved in 20 µL of TE for PCR reactions.

To prepare DNA representing input DNA, 200 µL of the chromatin solution (derived from 1 × 10^7 cells) was combined with 300 µL of elution buffer. Reverse of formaldehyde cross-links and isolation of the DNA was done as described for DNA eluted from the immunoprecipitates. DNA was resuspended in 20 µL of TE for PCR reactions. Two microliters of this solution corresponded to 20% of the input and other portions were diluted so that 2 µL contained 10%, 1%, or 0.1% of the input material. Two microliters of each immunoprecipitated DNA or input DNA were subjected to a PCR reaction in a final volume of 50 µL containing 10 pmoles of each primer, 200 µM dNTP, 2 or 4 µCi 32P-dCTP, 2.5 Units of Taq polymerase and buffer (Roche Diagnostics). Each sample was subjected to 20 cycles of: 94°C, 1 min; 60°C, 55 sec; 72°C, 50 sec; and followed by a final extension at 72°C for 10 min. PCR primers were the following: 70 ChIP A.1 spans from ~48 to ~24 and has the sequence 5’-AAAAAGCGCCGGAGATATAAATAA-3’; 70 ChIP A.2 spans from ~250 to +229 and has the sequence 5’-GCAGGCAATGTCTGGTAGTCT-3’; 70 ChIP B.1 spans from +1357 to +1374 and has the sequence 5’-CTGTGCGAGCCGCTATC-3’; 70 ChIP B.3 spans from +1730 to +1710 and has the sequence 5’-GCCCTTTGCCCGATCTAC-3’. Ten microliters of each PCR reaction was run on a 6% nondenaturing acrylamide gel. Gels were fixed in 10% acetic acid for 20 min and then dried. Radioactivity was detected in the dried gel using a phosphimager (Molecular Dynamics). The intensity of each band was quantified by volume analysis using ImageQuant software. The accuracy of the assay was assessed by determining how well the measurements for inputs matched the expected ratio of 0.1:1:10. In all cases, measurement with an expected value of 0.1 fell between 0.088 and 0.16, and the measurement with an expected value of 10 fell between 8.7 and 10.1.

Permanganate genomic footprinting

The fly line 17A10 containing a NELF-E RNAi construct controlled by the Gal4 UAS has been described previously (Enery et al. 2002). To direct expression of RNAi in salivary glands, the NELF-E RNAi flies were mated to the Bloomington stock #1824 [genotype: y1 w1; P[wre ^mwh+; = GawB[A1] that expresses Gal4 in salivary glands. Permanganate genomic footprinting was performed on isolated salivary glands as described previously (Weber et al. 1997; Tang et al. 2000). DNA breaks in the nontranscribed strand of the hsp70 genes at the 87A locus were detected using three overlapping primers that span the region from +165 to +229 [LM1–87A140, 5’-GATTTCAGTAGTTGCAGTTTT TAG-3’; LM2–87A140, 5’-TAGTTGCAGTGTATAGTTAGTAA TTACTTGG-3’; LM3–87A140, 5’-GCAGTTTTTAGTTAAT TACTCGGTGTTCG-3’].

Immunofluorescence analysis

Polytenes chromosome spreads were prepared as described by Champlin et al. (1991). Antibody against NELF-D was affinity-purified as described by Benyajati et al. (1997) and used at a dilution of 1/20. The antibody against the largest subunit of DSIF (dSpt5) was described by Kaplan et al. (2000), and it was used at a dilution of 1/150. The antibody against Pol IIa was monoclonal antibody 8WG16 obtained from Covance (Richmond CA), and it was used at a dilution of 1/10. All primary antibodies were detected with 1/250 dilutions of Alexa 488 or Alexa 568 conjugated secondary antibodies.

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NELF and DSIF cause promoter proximal pausing on the *hsp70* promoter in *Drosophila*

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