Transcriptional regulation of a yeast HSP70 gene by heat shock factor and an upstream repression site-binding factor

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SSA1, one of the heat-inducible HSP70 genes in the yeast Saccharomyces cerevisiae, displays a significant basal level of expression under optimal growth conditions. Although multiple sites related to the heat shock element (HSE) consensus sequences are present in the SSA1 promoter region, one of these, HSE2, plays a key role in basal expression. An upstream repression site (URS) located adjacent to HSE2 causes repression of basal expression but has little effect on heat-inducible expression of SSA1. A series of DNase I footprinting assays suggests that heat shock factor (HSF) and a URS-binding factor (URSF) can bind simultaneously to the adjacent binding sites HSE2 and URS under optimal growth conditions. URSF in extracts from heat-shocked cells does not bind (or binds very poorly) to the URS adjacent to HSE2. However, URSF in these extracts is able to bind the URS if the URS is separated from HSE2 or if the HSE is mutated such that HSF binding is abolished. These in vitro experiments are consistent with in vivo results showing that the URS is able to repress transcription driven by HSE2 both before and after heat shock if it is separated from HSE2. Our results are consistent with a model of repression in which URSF and HSF bind simultaneously to the adjacent binding sites URS and HSE2 prior to heat shock. After heat shock, however, binding of the two proteins to the adjacent sites is exclusive, perhaps due to modification of HSF known to occur upon heat shock. Because HSF binding predominates, repression by URSF is relieved upon heat shock.

[Key Words: Heat shock, negative regulation, upstream repression site-binding factor; heat shock factor; transcription]

Received February 15, 1991; revised version accepted April 3, 1991.

Expression of heat shock genes is induced upon temperature upshift or a variety of other environmental stresses. This gene activation is rapid and transient and, in most cases, appears to be regulated primarily at the transcriptional level (Bienz and Pelham 1987). Transcriptional activation of almost all heat shock genes in eukaryotes requires a highly conserved sequence element, referred to as a heat shock element (HSE). The consensus sequence of HSE was initially proposed to be the 14-bp palindromic sequence $\text{CnnGAAnnTTCnnG}$, where $n$ is any nucleotide (Pelham 1982; Pelham and Bienz 1982). Recently, it has been proposed that an HSE is described more accurately as a DNA element consisting of variable numbers of 5-bp units, $n$GAAn, in alternating orientation every 5 nucleotides (Amin et al. 1988; Xiao and Lis 1988). A positively acting transcription factor, called heat shock factor (HSF), which binds specifically to the HSEs, has been identified (Parker and Topol 1984; Wu 1984; Kingston et al. 1987; Sorger and Pelham 1987; Weiderrecht et al. 1987). More recently it has been shown that the 5-bp sequence $n$GAAn is the unit of interaction with HSF (Perisic et al. 1989). It has been proposed that HSF from the yeast Saccharomyces cerevisiae and Drosophila forms a trimer both in solution and when bound to DNA. Oligomerization seems to be essential for high-affinity binding of HSF to DNA (Perisic et al. 1989; Sorger and Nelson 1989).

Yeast HSF can bind to DNA prior to heat shock, whereas in human and Drosophila cells HSF is able to bind to DNA only after heat shock. HSF isolated from heat-shocked yeast cells is more highly phosphorylated than that from control cells, suggesting a modulation of HSF activity by phosphorylation (Sorger et al. 1987; Sorger and Pelham 1988). Thus, it appears that yeast HSF is not regulated at the level of DNA binding, rather, its capacity to activate transcription is regulated by heat shock. Yeast HSF has a low basal activity that is further induced upon heat shock. This is consistent with the fact that the gene encoding yeast HSF is essential at all growth temperatures (Sorger and Pelham 1988; Weiderrecht et al. 1988).

SSA1, one of the heat-inducible HSP70 genes in the yeast S. cerevisiae, also displays a significant basal level of expression under optimal growth conditions. The

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structure and regulation of the SSA1 promoter are quite complex. The upstream promoter region of SSA1 contains multiple sites closely related to the HSE consensus sequence and an upstream repression site [URS] [Slater and Craig 1987; Park and Craig 1989]. Although there are other HSEs in the SSA1 promoter, HSE2 and the URS play key roles in the control of basal expression of the SSA1 gene. A double point mutation in HSE2 abolishes >80% of the basal activity of the whole promoter. The URS, located to the 3' side and just adjacent to HSE2, represses the basal expression but has little effect on heat-inducible expression of the native SSA1 gene. It has been suggested that the basal expression of the SSA1 gene is under negative control mediated by a repressor [Park and Craig 1989].

To gain insights into the mechanism of repression by the URS, we performed a series of experiments, the results of which suggest that HSF and the URS-binding factor [URSF] can bind simultaneously to the adjacent binding sites prior to heat shock. However, in heat-shocked cells, the two proteins appear to be unable to bind to the wild-type SSA1 promoter simultaneously; and because HSF binding predominates, repression is relieved.

Results

Identification of a protein (URSF) that binds to the URS region by DNase I footprinting and methylation protection experiments

Previous analyses of deletion and point mutants indicated that an upstream repression site is located adjacent to HSE2, on its 3' side, in the SSA1 promoter. By gel-shift assays using a DNA fragment carrying only the URS (from −169 to −194 with respect to transcription start site) we were able to detect a URS-specific binding activity in yeast whole-cell extracts [Park and Craig 1989]. To gain insight into the mechanism of repression by the URS, we performed a series of DNase I footprinting analyses using a DNA fragment of the SSA1 promoter (from −261 to −131) containing both HSE2 and the URS. Whole-cell extracts were prepared from yeast cells grown at 23°C, and the HSE- and URS-binding activities were enriched by chromatography on a heparin–Sepharose column. The DNase I protection pattern of the top strand of the wild-type DNA fragment is shown in Figure 1A. The protected region extended from −170 (−174) to −206 (−204), which includes both HSE2 and the URS (numbers in parentheses indicate uncertainties in the footprint boundaries due to partial protection or to poor cleavage of the DNA at those regions). To determine whether this protection was due to HSF and/or a URS-binding protein, we performed DNase I footprinting analyses using DNA fragments carrying mutations in either HSE2 or the URS [see Fig. 1B], which have been shown previously to destroy either HSE2 or the URS activity in vivo [Park and Craig 1989]. The protected region of the top strand of the URS mutant probe extended from −182 to −206, presumably due to binding of HSF. In contrast, the protected region of the HSE2 mutant probe extended from −170 to −187, indicating that protection of the URS region is the result of the binding of a distinct URS-binding factor, which we call URSF.

We also observed regions protected from DNase I cleavage both 5' and 3' of HSE2 and the URS (Fig. 1A). Because deletions removing these sites did not significantly affect either the basal or heat-inducible expression [Park and Craig 1989], we did not consider proteins binding to these sites in this study. They may be involved in regulation of SSA1 under other conditions.

Because the DNA-binding sites of HSF and URSF determined by DNase I footprinting were not separated from each other, we wanted to gain more detailed insight into these protein–DNA interactions. To determine specific nucleotide contacts of each protein, we performed methylation protection experiments using the wild-type DNA fragments and the same DNA fragments carrying either the HSE2 or URS mutation used in the DNase I footprinting. As shown in Figure 2A, strong protection at one G [−196] and weaker protection at four G residues [−184, −183, −181, and −180] were observed with the wild-type top strand. With a probe carrying a mutation in HSE2, the four G residues mentioned above were still protected, with a probe carrying a mutation in the URS, protection at only one G [−196] was observed. A strong protection was observed at two G residues [−199 and −189] on the wild-type bottom strand [Fig. 2B]; no G residues were protected on the bottom strand of the HSE2 mutant probe [data not shown]. The results indicate that a G residue [−196] of the top strand and two G residues [−199 and −189] of the bottom strand, all of which occur in the HSE consensus sequence, make contact with HSF. Four G residues [−184, −183, −181, and −180] were protected by URSF. We also observed some weaker protection of G residues at the 3' boundary and outside of the region protected by URSF as determined by DNase I footprinting. It is not clear at this point whether this protection is due to some DNA structural change caused by URSF binding or a protein binding at the 3'-flanking region of the URS.

Initially, proteins were prepared for DNase I footprinting by loading onto the heparin–Sepharose column at 0.1 M KCl and then eluting with successive steps of 0.3 and 0.6 M KCl. Most of the HSE- and URS-binding activities were found in the 0.6 M KCl fraction. Subsequently, we found that the URS- and the HSE-binding activities were partially separated by adding a 0.4 M KCl elution step in the heparin–Sepharose column fractionation. The URS-binding activity was found in the 0.6 M KCl fraction from the column, whereas most of the HSE-binding activity was found in the 0.4 M KCl fraction. The regions protected from DNase I digestion by each fraction were partially overlapping. The URSF-containing fraction protected from −187 to −174 (−170) of the top strand [Fig. 3A] and −189 (−193) to −178 (−174) of the bottom strand from DNase I cleavage [Fig. 3B]. HSF protected from −206 (−204) to −182 of the top strand [Fig. 3A] and from −208 to −183 of the bottom strand [Fig. 3B].
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This separation of binding activity also supports the idea that different proteins bind to the HSE and URS. A summary of the DNase I footprinting and methylation protection results is shown in Figure 4.

Identification of URSF by UV cross-linking

To identify URSF directly we performed UV cross-linking experiments using a DNA fragment (from –261 to –131) carrying both HSE2 and the URS as a probe. A protein migrating as a 40-kD polypeptide was detected in the 0.4–0.6 M KCl fraction that is enriched for URSF [Fig. 5]. When the same DNA fragment carrying a point mutation in the URS (URS–M2; see Fig. 1B) was used as a probe, this band was not apparent, indicating that this protein was specifically interacting with the URS. Detection of the major 40-kD polypeptide was dependent on the addition of protein and UV radiation. More extensive nuclease digestion of the UV-irradiated binding mixtures did not alter the mobility of this major band. Because the covalent attachment of short oligonucleotides to proteins has only minor effects on the mobility of the proteins in SDS gel, the results indicate that a protein of Mr 40,000 binds specifically to the URS.

As an alternative means of identifying URSF, proteins were separated by SDS-PAGE, transferred to nitrocellulose, and probed with a 32P-labeled 32-bp oligonucleotide carrying the URS. A single band of similar size to that obtained by UV cross-linking was detected using extracts prepared from control and heat-shocked cells [data not shown].

We have detected, by DNase I footprinting, other proteins interacting with sites flanking HSE2 and the URS [see Fig. 1A]. A major band of 100 kD observed by UV cross-linking with both wild-type and the URS–M2 mutant probe (Fig. 5) could be due to a protein bound to one of these flanking sites, or, alternatively, to nonspecific binding. The protein migrating at 150 kD and giving a weak signal [Fig. 5] is presumably HSF. We observed a stronger band of the same relative molecular mass when the 0.3–0.4 M KCl fraction containing most of the HSE-binding activity was used, and the 150-kD size is consistent with the apparent relative molecular mass of HSF published previously by another group [Sorger and Pelham 1987].

Figure 1. (A) DNase I protection of the wild-type and mutant SSA1 promoters by control extract. An XhoI–PvuI fragment from the plasmid pZHSE2-137A [see Materials and methods], which carries the SSA1 upstream sequences from –261 to –131, was labeled at the 3’ end, incubated for 10 min at room temperature without protein (−) or with 150 µg of the 0.3–0.6 M KC1 fraction (+) from a heparin–Sepharose column, and partially digested with DNase I as described in Materials and methods. (WT) The wild-type SSA1 DNA fragment; (URS–M2) the same DNA fragment containing a point mutation in the URS; (HSE2–M) the same DNA fragment carrying a mutation in HSE2. (Lane G + A) G + A chemical sequencing reaction of the wild-type probe. Solid lines indicate the span of the footprint; dotted lines connected to the solid lines indicate uncertainties in the footprint boundaries due to poor cleavage of the DNA at those regions or to partial protection. Dashed lines indicate the regions where the protection is lost due to mutations. DNase I protection pattern of the wild-type bottom strand is shown in Fig. 7B. The sequence of the protected region is shown in Fig. 4. (B) Schematic representation of the SSA1 promoter and its mutant derivatives. Each diamond represents an HSE–homologous site [Slater and Craig 1987]. The enlarged region shows the sequence of HSE2 and the URS region. (+1) The transcription start site. The long arrowheads mark the component GAA repeats with relative spacing, which has been proposed to comprise a functional HSE [Amin et al. 1988; Xiao and Lis 1988]. The box encompasses the consensus sequence of the URS [Luche et al. 1990]. The base substitutions in HSE2 (HSE2–M) and in the URS (URS–M1, URS–M2) are shown below the wild-type (WT) sequence.
Figure 2. Methylation protection patterns of HSE2 and the URS. DNA methylation protection patterns of SSA1 DNA fragments carrying the wild type (WT), mutations in HSE2 (HSE2-M), or the URS (URS-M2) are shown. [A] Top strand; [B] bottom strand. Positions of methylated G residues that were strongly or weakly protected by the factors are marked by solid or open triangles, respectively. (Lanes G + A) G + A chemical sequencing reactions of 3' the wild-type probe. Bold letters in the sequence indicate the mutated residues.

URS can repress transcription even after heat shock when it is separated from HSE2

We showed previously that the URS represses the basal expression of the SSA1 gene but has little effect on heat-inducible expression. In contrast, the URS can repress transcription of a heterologous promoter both before and after heat shock (Park and Craig 1989; see also Fig. 6A). The 137-bp DNA fragment of the SSA1 promoter containing both HSE2 and the URS (from -261 to -131) was inserted downstream of the CYC1 upstream activating sequence (UAS) in the CYC1-lacZ fusion vector. Insertion of the 137-bp DNA fragment carrying a mutation in HSE2 but a wild-type URS (HSE2-M) almost completely abolished CYC1 UAS activity both at 23°C and after shift to 37°C. In contrast, insertion of the 137-bp fragment carrying a mutation in the URS (URS-M1 or URS-M2) did not result in repression of CYC1 transcription [Fig. 6A].

Because the URS could repress expression of a heterologous promoter both before and after heat shock, we decided to test whether the URS could repress transcription driven by HSE2 even after heat shock if it is separated from HSE2. The 137-bp fragment carrying a mutation in HSE2 (HSE2-M) but a wild-type URS was inserted 3' of HSE2 in the CYC1-lacZ fusion vector, which carries HSE2 in place of the CYC1 UAS. As shown in Figure 6B, CYC1 transcription driven by HSE2 was re-

Figure 4. Summary of sequences of HSE2 and the URS region that are protected from DNase I cleavage and methylation by HSF and URSF. Long arrowheads represent repeated GAA blocks within the 5-nucleotide heat shock regulatory module (Amin et al. 1988; Xiao and Lis 1988). Solid lines indicate the span of DNase I footprint; dashed lines indicate uncertainties in the footprint boundaries due to poor cleavage of the DNA at those regions or due to partial protection. Positions of methylated G residues that were protected by HSF or URSF are marked by small solid or open triangles, respectively.
Figure 5. Identification of UR SF by UV cross-linking. A wild-type SSA1 promoter fragment (from -261 to -131) [lanes 1-3] or its derivative carrying a mutation in the URS (lane 4) was incubated with 60 μg of the 0.4-0.6 M KCl fraction from heparin-Sepharose chromatography. The binding reaction mixtures were UV-irradiated as described in Materials and methods, and the mixture was separated on a polyacrylamide gel. (Lane 1) No protein added; (lane 2) UV irradiation omitted; (lane 3) wild-type probe plus extracts; (lane 4) URS mutant [URS-M2] probe plus extracts. The positions of protein molecular mass markers are indicated at left.

pressed ~18-fold after heat shock when the URS was separated from HSE2. Although the repression ratio is smaller than that of basal activity, this result indicates that the URS can repress transcription driven by HSE2 after heat shock when it is separated from HSE2. As expected, insertion of the 137-bp fragment carrying a mutation in the URS [URS-M1 or URS-M2] did not result in repression. β-Galactosidase activities of control and heat-shocked cells carrying these plasmids were elevated compared to the parental plasmid [pZJHSE2-26], probably due to the presence of two copies of HSE in the constructions. Taken together, these results suggest a heat shock-independent mode of repression by the URS if it is separated from HSE2.

Binding of UR SF to the URS of the native SSA1 promoter is greatly reduced after heat shock

To test binding of UR SF to the URS in the native SSA1 promoter after heat shock, we performed DNase I footprinting using extracts prepared from heat-shocked cells. Whole-cell extracts were prepared from yeast cells grown at 23°C and shifted to 37°C for 30 min prior to harvest, and then fractionated on a heparin–Sepharose column as described above. The 0.3–0.6 M KCl fraction protected from -206 to -182 of the top strand [Fig. 7A] and from -208 to -183 of the bottom strand [Fig. 7B] of the wild-type SSA1 promoter fragment [cf. lanes HS and C]. The region protected with heat shock extract from DNase I digestion includes HSE2 and only the 5' portion of the URS, suggesting that UR SF prepared from heat-shocked cells is not able to bind [or binds very poorly] to the native SSA1 promoter. The result is consistent with the fact that, in vivo, the URS represses the basal expression of the SSA1 gene but has little effect on heat-inducible expression [Park and Craig 1989].

However, it was shown by β-galactosidase assays that the URS can repress transcription driven by HSE2 even after heat shock when it is separated from HSE2 [see above]. Thus, we decided to test whether UR SF in extracts from heat-shocked cells can bind to the URS by a gel-shift assay. Previously, we showed by gel-shift assays that a protein specifically interacts with the URS. The binding could be competed by addition of excess unlabeled oligonucleotide carrying the URS but not oligonu-
Figure 7. DNase I protection pattern of the SSA1 promoter fragment by control and heat shock extracts. The SSA1 promoter fragment (from −261 to −131) was used as a probe (see legend to Fig. 1A). Whole-cell extracts prepared from cells grown at 23°C (C) or shifted to 37°C for 30 min prior to harvest (HS) were fractionated on a heparin-Sepharose column as described in Materials and methods. The 0.3–0.6 M KCl fraction was incubated for 10 min at room temperature with extract from heat-shocked (HS) or control (C) cells, or without protein (−). [G + A] G + A chemical sequencing reaction of the probe.

Discussion
URSF, a repressor potentially regulating a variety of yeast promoters

Using DNase I footprinting and UV cross-linking, we have identified a factor that binds to the URS located very near to HSE2 of the HSP70 gene, SSA1. This URSF has an apparent Mr of 40,000. In crude extracts, URSF from control cells, but not heat-shocked cells, binds to the URS of the wild-type SSA1 promoter; a single point mutation in the URS diminishes binding of URSF to the site. These results are consistent with effects of the URS on in vivo expression: repression under optimal growth conditions but little effect on expression of SSA1 after heat shock. Therefore, the data strongly suggest that URSF is a repressor regulating the SSA1 gene.

The URS in the SSA1 gene shares strong homology with URS1 in the CAR1 gene. It has been suggested that the CAR1 URS consists of a 9-bp sequence, AGCCGCGA, based on saturation mutagenesis results (Luche et al. 1990). Our DNase I footprinting assays show that the sequence with one mismatch to the CAR1 URS, in the inverted orientation, is located in the middle of a region protected from DNase I cleavage by heat shock extract. In contrast, only the HSE2 region was protected with the wild-type probe. Protection of the URS region of the HSE2 mutant probe was indistinguishable when either control or heat shock extracts were used (see Figs. 1A and 8C). Together, the results suggest that URSF cannot bind to the URS in the wild-type SSA1 promoter after heat shock because of the binding of HSF nearby.
Negative regulation of a heat shock gene

been shown that the sequence is contained in a region associated with negative control of gene expression (Cohen et al. 1987; Sumrada and Cooper 1987). Luche et al. (1990) also reported a protein interacting with the CAR1 URS by gel-shift assays. Competition studies using DNA fragments from the SSA1 promoter indicate that the protein interacting with the CAR1 URS may be the same as the URSF that regulates SSA1. It is therefore possible that the URSF studied here also represses transcription of a variety of yeast genes. However, a family of related proteins might share DNA-binding specificity, but interact specifically with different transcription factors.

Repression of basal transcription of SSA1 by the URS

Two basic mechanisms of repression of transcriptional initiation have been considered. The most commonly observed repression mechanism in prokaryotes involves competition between a repressor protein and general transcription factors such as RNA polymerase for binding to closely placed sites. A related form of competition found in eukaryotes involves competition between a repressor and an activator protein that bind overlapping or closely located sites. In the second type of mechanism, an activator and a repressor bind to nonoverlapping DNA sequences. Protein–protein interactions between the two proteins may prevent the activator from making productive interactions with the general transcription machinery (Levine and Manley 1989).

We performed a series of DNase I footprinting assays with increasing amounts of a fraction containing both HSF and URSF. If the two factors cannot bind at the same time and there is competition between HSF and URSF for binding to the adjacent sites, one would expect to see a partial protection in both the HSE2 and the URS region. However, the HSE2 region became fully saturated even at low protein concentrations, at which point the URS region was only partially protected by URSF (data not shown). A further increase in the amount of total protein resulted in increased protection of the URS region but did not affect protection of HSE2 region by HSF (see Figs. 1A and 7A and B). Thus, DNase I footprinting analyses using control [non-heat shock] extracts indicate that URSF and HSF can bind simultaneously to the HSE2–URS region of the SSA1 promoter. The regions protected by HSF and URSF partially overlap. However, such overlap does not preclude the possibility that both factors bind to the adjacent sites simultaneously, because DNase I may be unable to cleave nucleotides very close to the site of contact of HSF and URSF with the DNA. Methylation protection experiments indicate that

Figure 8. Interaction of URSF from heat-shocked cells with the URS on DNA fragments lacking or having altered HSE. (A) Detection of URSF in the control and heat shock extracts. A gel-shift assay was performed with 32P-labeled 32-bp double-stranded oligonucleotide carrying the URS (see Materials and methods) and 30 µg of whole-cell extract prepared from cells grown at 23°C (C) or heat-shocked cells (HS). F and B indicate free DNA and protein-bound DNA, respectively. (B) URSF in extracts from heat-shocked cells can bind to a URS separated from HSE2 even after heat shock. An XhoI–RsaI fragment from plasmid pH104 (see Materials and methods), which carries the wild-type HSE2 separated from the URS by 73 bp, was labeled at the 3’ end and incubated without protein (−) or with 100 µg of 0.3–0.6 M KCl fraction (+) prepared from heat-shocked cells. (C) The URS of the SSA1 promoter is protected by URSF from heat-shocked cells when the adjacent HSE2 is mutated. The wild-type SSA1 promoter fragment (from −261 to −131) [WT] or a fragment carrying mutations in HSE2 [HSE2–M] (see Fig. 1B) was incubated without protein (−) or with the 0.3–0.6 M fraction (+) prepared from heat-shocked cells and partially digested with DNase I. Equal amounts of protein (~100 µg) were used for both lanes WT and HSE2–M.
HSF and URSF do not contact the same nucleotides. We were unable to resolve a guanine HSF–URSF–DNA ternary complex from an HSF–DNA complex by gel-shift assays, perhaps due to the very small size of URSF compared to HSF. HSF appears to bind to DNA as a trimer ([Sorger and Nelson 1989]), which would contribute ~450 kD. Our UV cross-linking data suggest that URSF is a protein of only ~40 kD.

In conclusion, repression of basal expression of SSA1 does not appear to be caused by URSF preventing the activator HSF from binding to the nearby HSE. It is more likely that URSF inhibits basal transcription by interfering with the function of HSF bound at HSE2, perhaps by preventing proper interaction between HSF and the basic transcription machinery. Alternatively, URSF might inhibit transcription without interacting with HSF, perhaps by affecting looping of the DNA. The data available at this time do not allow us to distinguish between these and other mechanisms of repression.

A model for the lack of repression by URSF after heat shock
Repression of expression of the SSA1 gene by the URS was seen only prior to heat shock. URSF in extracts prepared from heat-shocked cells binds very poorly to the URS of the wild-type SSA1 promoter as assayed by DNase I footprinting. However, we have shown in three different ways that the DNA-binding ability of URSF is unchanged upon heat shock. First, URS-binding activity was detected in both control and heat shock extracts by a gel-shift assay using an oligonucleotide containing the URS. Second, HSF and URSF in heat shock extracts can bind to their own binding sites when the two sites are separated. Third, DNase I footprinting analyses using an SSA1 promoter DNA carrying an HSE2 mutation showed that URSF can bind to the URS even after heat shock when HSF is no longer able to bind to HSE2. URSF can also repress transcription of a heterologous promoter in vivo after heat shock when the URS is not next to a wild-type HSE2. These data are consistent with the idea that there is no apparent change in the DNA-binding ability of URSF upon heat shock and URSF cannot bind to the URS in the wild-type SSA1 promoter after heat shock due to the binding of HSF to HSE2 nearby.

While the activity of HSF is greatly increased upon heat shock, it has been reported that the affinity of HSF for DNA does not change ([Sorger et al. 1987]). We confirmed this result by a series of gel-shift assays [data not shown]. Thus, it is unlikely that the poor binding of URSF to the wild-type SSA1 promoter is due to an increase in HSF binding after heat shock. However, from footprinting assays, it appears that either HSF is more abundant or has a higher affinity for its binding site than that of URSF for the URS ([H.-O. Park and E.A. Craig, unpubl.]). Because complete protection of HSE can be achieved at much lower protein concentrations. Thus, if HSF and URSF cannot bind simultaneously after heat shock, competition for binding between HSF and URSF could result in HSF being the predominant protein bound to the HSE2–URS region.

Although URSF and HSF in control cell extracts were partially separated by heparin–Sepharose column fractionation, HSF in heat-shock extracts cofractionated with URSF ([H.-O. Park and E.A. Craig, unpubl.]), this might be caused by modification of HSF upon heat shock. It has been shown that HSF isolated from heat-shocked yeast cells is more highly phosphorylated than that isolated from control cells ([Sorger and Pelham 1988]). We have observed by immunoblotting that HSF in our extracts prepared from heat-shocked cells migrates more slowly on SDS gels than that from cells grown at 23°C [data not shown], consistent with the altered migration caused by phosphorylation reported previously. Thus, it is tempting to speculate that phosphorylation of HSF is responsible for release from the URS-mediated repression after heat shock, and this coupled with the activation of HSF causes a dramatic increase in SSA1 expression upon heat shock.

We propose the following model, which accommodates our data: Both the control form of HSF and URSF can bind to the adjacent binding sites, HSE2 and URS, prior to heat shock. URSF represses transcription by interfering with the ability of the control form of HSF to activate transcription. After heat shock, modified HSF bound to the HSE2 region does not allow simultaneous binding of the two proteins to the adjacent sites, perhaps due to steric hindrance caused by a conformational change of HSF or charge repulsion due to phosphorylation of HSF that occurs after a heat shock (Fig. 9). Puriﬁcation of URSF and analysis of possible interactions between HSF and URSF will allow us to understand the mechanism of repression by the URS in more detail.

Materials and methods
Yeast stains and plasmids
Strain T69 [MATa ura3-52 lys2-801 ade2-101 Δtrp1] was used as a host for all plasmids for β-galactosidase assays. Protease-defi...
cient strain BJ2168 [MATa prb1-1122 pep4-3 prc1-407 leu2 trp1 ura3-52 gal2] was used for extract preparation.

Plasmids pZHSE2-26 and pZL4 (Fig. 6) were described previously (Slater and Craig 1987). All derivatives of these plasmids were generated by inserting a 137-bp XhoI fragment (containing an SSA1 promoter fragment from -261 to -131) carrying either a wild-type or mutated HSE2, or URS. The mutations in HSE2 and the URS have been described previously [Park and Craig 1989] and are also shown in Figure 1B.

Plasmids pZHSE2-137A and pZHSE2-137B, used to generate probes for DNase I footprinting assays (see Fig. 1A legend), were constructed by XhoI partial digestion of pZHSE2-137 [Slater and Craig 1987] and filled in with Klenow, generating a Prot site. Thus, pZHSE2-137A or pZHSE2-137B contains either a downstream or an upstream XhoI site, respectively.

Plasmid pHPl04, used to generate a probe used for DNase I footprinting shown in Figure 8B, carries the 137-bp XhoI fragment with the HSE2 mutation [HSE2-M] at the 3′ end of the 26-mer of HSE2.

Enzyme assays

Yeast cells were transformed with each plasmid and grown on glucose-based synthetic complete media lacking uracil. β-Galactosidase activity in yeast cells was determined before (23°C) and after (37°C) heat shock during logarithmic growth as described previously [Park and Craig 1989].

Extract preparation and heparin–Sepharose column chromatography

Yeast cells grown rich, glucose-based media to an OD₆₀₀ of 1.3–1.5 at 23°C were used for preparation of control extracts, as described [Park and Craig 1989]. For preparation of heat shock extracts, cells grown to an OD₆₀₀ of 0.9–1.0 at 23°C were shifted to 37°C for 30 min prior to harvest. Yeast whole-cell extracts were prepared as described previously [Park and Craig 1989]. Whole-cell extracts were loaded directly on a heparin–Sepharose column and washed first with three column volumes of buffer A [20 mM HEPES at pH 8.0, 1 mM MgCl₂, 0.1 mM EDTA, 0.5 mM PMSF, 0.2 mM DTT] containing 0.1 mM KCl, and then three column volumes of buffer A containing 0.3 M KCl. Both HSE- and URS-binding activities were eluted with two column volumes of buffer A containing 0.6 M KCl. The 0.3–0.6 M fraction was precipitated with ammonium sulfate, [final 50% saturation] or concentrated using a Centricon-10 concentrator [Amicon, Danvers, MA], and dialyzed overnight against buffer A containing 0.1 M KCl. Following dialysis, insoluble materials were precipitated by centrifugation and the supernatant was stored frozen at -70°C. Protein concentrations were determined by the Bradford method [Bradford 1976] using reagents from Bio-Rad.

Gel-shift assays, DNase I footprinting, and methylation protection analyses

Labeling of DNA fragments for probes and the binding reactions was carried out as described previously [Park and Craig 1989]. The 32-bp oligonucleotide, described previously [Park and Craig 1989] and used as a probe for gel-shift assays, is shown below:

\[5'-TCGACACGTTCCATCGGCGGCAAAAGGGAGAC-3']

5′-GTGCAAGTGACGCCGCAAAGGAGAC-3′

A 137-bp XhoI-PvuI fragment from plasmid pZHSE2-137A (top strand) or pZHSE2-137B (bottom strand) was used for DNase I footprinting assays. The DNA fragment was labeled at the 3′ end with [α-32P]dCTP and Klenow fragment. For DNase I footprinting analysis shown in Figure 8B, a 283-bp XhoI-Rsal fragment from plasmid pHPl04 was generated by partial XhoI digestion and labeled at the XhoI end. For each binding reaction for DNase I footprinting assays, 0.2–0.5 ng of 32P-labeled DNA fragment was incubated with 3 μg of poly[d(I-C)/d(I-C)] and 80–150 μg of the 0.3–0.6 M KC1 fraction from a heparin–Sepharose column. The binding reaction was carried out at room temperature for 5–10 min, followed by DNase I digestion usually at room temperature for 1 min. The amount of DNase I [Sigma] was determined empirically and ranged from 20 to 400 ng. After DNase I digestion, the mixture was extracted with phenol/chloroform, precipitated with ethanol, and subjected to electrophoresis on a 7 M urea sequencing gel.

For methylation protection analysis, the binding reactions were carried out as described above, followed by partial methylation at the guanine residues, as detailed by Maxam and Gilbert (1977).

UV cross-linking

Both the wild-type and the URS mutant DNA fragments [carrying the SSA1 promoter sequences from -131 to -261] were used as probes for UV cross-linking experiments, which were performed by the method of Chodosh et al. [1986] with slight modifications. The DNA was labeled using Klenow fragment and random primer in the presence of 50 μM dATP, dGTP, 5-bromo-2′-deoxyuridine triphosphate [Pharmacia], and 5 μM [α-32P]dCTP. Protein samples of the 0.4–0.6 M fraction from a heparin–Sepharose column were incubated with 1 ng of uniformly labeled DNA and 3 μg of poly[d(I-C)/d(I-C)] in 30 μl of binding buffer at room temperature. The mixture in an Eppendorf tube was irradiated under a Fotedye UV lamp (maximum emission wavelength, 310 nm; maximum intensity, 7000 mW/cm²) at a distance of 4.5 cm from the UV source. The mixture was brought up to 10 mM CaCl₂, and DNA was digested for 30 min at 37°C with 3 μg of DNase I [Sigma] and 1 unit of micrococcal nuclease [Pharmacia]. The mixture was then electrophoresed on an SDS–7.5% polyacrylamide gel, and the gel was dried onto 3M Whatman filter paper and autoradiographed. The positions of protein molecular mass markers were determined by prestained protein markers from Sigma.

Acknowledgments

We are grateful to Peggy Famham and Charles Nicolet for critical reading of the manuscript. We also thank Peggy Famham, Carol Gross, Charles Nicolet, Anna Means, Pil Jung Kang, and Mike Young for helpful discussions during the progress of this work. This work was supported by a U.S. Public Health Service grant from the National Institutes of Health to E.A.C. and by the Lucille P. Markey Charitable Trust, Miami, Florida to H.-O.P.

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*Genes Dev.* 1991, 5:
Access the most recent version at doi:10.1101/gad.5.7.1299

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