Deciphering the role of the signal- and Sty1 kinase-dependent phosphorylation of the stress-responsive transcription factor Atf1 on gene activation

Received for publication, May 2, 2017, and in revised form, June 23, 2017. Published, Papers in Press, June 26, 2017. DOI 10.1074/jbc.M117.794339

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Edited by Joel Gottesfeld

Adaptation to stress triggers the most dramatic shift in gene expression in fission yeast (Schizosaccharomyces pombe), and this response is driven by signaling via the MAPK Sty1. Upon activation, Sty1 accumulates in the nucleus and stimulates expression of hundreds of genes via the nuclear transcription factor Atf1, including expression of atf1 itself. However, the role of stress-induced, Sty1-mediated Atf1 phosphorylation in transcriptional activation is unclear. To this end, we expressed Atf1 phosphorylation mutants from a constitutive promoter to uncouple Atf1 activity from endogenous, stress-activated Atf1 expression. We found that cells expressing a nonphosphorylatable Atf1 variant are sensitive to oxidative stress because of impaired transcription of a subset of stress genes whose expression is also controlled by another transcription factor, Pap1. Furthermore, cells expressing a phospho-mimicking Atf1 mutant display enhanced stress resistance, and although expression of the Pap1-dependent genes still relied on stress induction, another subset of stress-responsive genes was constitutively expressed in these cells. We also observed that, in cells expressing the phospho-mimicking Atf1 mutant, the presence of Sty1 was completely dispensable, with all stress defects of Sty1-deficient cells being suppressed by expression of the Atf1 mutant. We further demonstrated that Sty1-mediated Atf1 phosphorylation does not stimulate binding of Atf1 to DNA but, rather, establishes a platform of interactions with the basal transcriptional machinery to facilitate transcription initiation. In summary, our results provide evidence that Atf1 phosphorylation by the MAPK Sty1 is required for oxidative stress responses in fission yeast cells by promoting transcription initiation.

Several mitogen-activated protein (MAP) kinase pathways allow eukaryotic organisms to respond to environmental challenges by triggering stress-dependent gene expression programs. Upon exposure to signals, phosphorylated MAP kinase accumulates in the nucleus and triggers phosphorylation of transcription factors (TFs). The activated TFs are then able to translate extracellular cues into specific cellular responses by adapting the complex RNA polymerase II (Pol II) transcriptional machinery into particular sets of genes and mediate specific changes in the gene expression program (1). The fission yeast Schizosaccharomyces pombe responds to environmental stressors by inducing a complex signal transduction pathway meant to allow survival: the Sty1/Spc1 MAP kinase pathway. The pathway is induced by many stress conditions and triggers a wide transcriptional shift of the gene expression program (2, 3). A common consequence of the different signals is activation by phosphorylation of the MAP kinase Sty1 (4, 5). Then, Sty1 transiently accumulates in the nucleus, where it promotes transcriptional activation or repression of genes in an at least partially Atf1-dependent manner (4, 6–8). Atf1 is a basic zipper (bZIP)-containing TF that heterodimerizes with another bZIP protein called Pcr1; even though the phenotypes of cells lacking one or another TF are not identical, they have overlapping functions (9, 10), and Atf1 seems to be the direct substrate of the MAP kinase Sty1 (8). Sty1 phosphorylation is required to trigger both nuclear accumulation as well as activation of its kinase activity because constitutive nuclear accumulation of the kinase is not sufficient to induce phosphorylation of its main substrate, Atf1, or to activate transcription (9, 11). In response to toxic but not lethal extracellular hydrogen peroxide (H2O2), more than 500 genes are up-regulated more than 2-fold. Their induction depends on Sty1 and, to a lesser extent, on Atf1 (2, 3). Even though a lot of work has been done to identify and characterize the downstream effectors of activated Sty1–Atf1 on transcription regulation, such as the SAGA complex (12), the main role of the Sty1 kinase activity on Pol II-dependent transcription of stress genes is not clear. A first possibility, based on the ortholog kinase HOG1 of Saccharomyces cerevisiae (for a review, see Ref. 13), is

This work was supported by the Ministerio de Economía y Competitividad (Spain), PLAN E and FEDER (BFU2015-68350-P to E. H.), and by 2014-SGR-154 from Generalitat de Catalunya (Spain) (to E. H.). The authors declare that they have no conflicts of interest with the contents of this article. This article contains supplemental Figs. S1–S4 and Tables S1 and S2. The data discussed in this publication have been deposited in the NCBI Gene Expression Omnibus and are accessible through GEO series accession number GSE97057.

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‡ Recipient of a Maria de Maeztu predoctoral fellowship from the Ministerio de Economía y Competitividad (Spain).
§ Recipient of a Juan de la Cierva postdoctoral contract from the Ministerio de Economía y Competitividad (Spain).
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The abbreviations used are: MAP, mitogen-activated protein; TF, transcription factor; Pol II, RNA polymerase II; bZIP, basic zipper; MM, minimal medium; YE, rich medium; emp, empty vector; ORFs, open reading frames.
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that the kinase participates directly in transcription initiation and/or even elongation; in this model, the MAP kinase Sty1 would only use Atf1 as an anchor to the stress promoters (10, 14, 15), where activated Sty1 could regulate the access or activity of the transcriptional machinery through phosphorylation of unknown substrate(s). A second proposal is that nuclear and activated Sty1 enhances, by phosphorylating Atf1, the affinity of the TF for its cAMP response element sites at stress promoters: only phosphorylated Atf1 would display low dissociation constants toward the cAMP response element sites. Although a genome-wide study using ChIP sequencing of total immunoprecipitated Atf1 indicates that there is a significant enhancement of TF binding to DNA upon activation by Sty1 (16), ChIP of individual stress genes reported very modest recruitment, if any, of Atf1 to specific stress promoters upon stress imposition (9, 15). Finally, it has also been proposed that the main role of Sty1 in promoting Atf1 function is by inhibiting its ubiquitin-independent degradation (10, 17) so that stabilization and accumulation of Atf1 upon phosphorylation would be the gene activation–triggering event; in this case, enhanced TF concentration would increase promoter occupancy.

Another important aspect of this regulatory cascade is that the atf1 gene is also up-regulated upon stress as part of a feedback loop. Thus, oxidative stress triggers transcription of this gene 4-fold (2, 3), which may contribute to enhancing Atf1 protein levels after stress. Therefore, we decided to perform a characterization of the role of Sty1-dependent Atf1 phosphorylation by expressing, from a constitutive promoter, HA-tagged wild-type Atf1 and several mutants lacking some of the 11 S/TP phosphosites. We demonstrate here that the transcriptional profile of ∆atf1 cells expressing HA-Atf1 is very similar to that of wild-type cells despite the fact that the concentration of the TF is constant. Cells expressing Atf1 mutants lacking the S/TP sites are unable to trigger transcription of a subset of stress genes, whereas Atf1 mutants with substitutions mimicking phosphorylation are able to trigger constitutive or inducible transcription of the stress genes in a Sty1-independent manner.

Results

Expression of Atf1 from a constitutive promoter does not alter the pattern of activation of stress genes

The atf1 mRNA is up-regulated four-times upon H₂O₂ exposure in an Atf1-dependent manner (2). We expressed HA-Atf1 under the control of the constitutive promoter sty1; the gene chimera, coding for HA-Atf1 containing the 11 MAP kinase phosphorylation sites (Fig. 1A), was integrated at the leu1 locus of cells lacking Atf1. Expression of the HA-Atf1 chimera suppresses the sensitivity to peroxides of cells deficient in Atf1 on solid plates (Fig. 1B). As shown by Western blotting, a shift in electrophoretic mobility in extracts from stressed cells identical to the wild-type protein, whereas the HA-Atf1.10M and HA-Atf1.11M (data not shown) proteins did not seem to have a significant change in mobility. Concomitantly, although the HA-Atf1.1M mutant was fully able to suppress the sensitivity to peroxides of strain ∆atf1, expression of the HA-Atf1.10M and 11M mutants did not alleviate this phenotype (supplemental Fig. S1C). We then constructed a chimeric gene coding for HA-Atf1.10D to render a phospho-mimicking mutant (serine or threonine sites were mutated to alanine or isoleucine to avoid phosphorylation). As shown by Western blotting in supplemental Fig. S1B, the HA-Atf1.1M mutant displays an apparent shift in electrophoretic mobility in extracts from stressed cells identical to the wild-type protein, whereas the HA-Atf1.10M and HA-Atf1.11M (data not shown) proteins did not display any detectable mobility shift upon stress (Fig. 1F). Concomitantly, although expression of HA-Atf1.10M was not able to suppress the sensitivity to peroxides of strain ∆atf1 (supplemental Fig. S1C and Fig. 1G), expression of HA-Atf1.10D alleviated this phenotype (Fig. 1G).

We introduced lower number of substitutions in Atf1 to try to determine exactly which modification/s was/were essential for the role of Atf1 in oxidative stress survival. As shown in supplemental Fig. S2, strain ∆atf1 expressing the mutant named HA-Atf1.6M, lacking sites 5 to 10 in Atf1, was as sensitive to growth on peroxide-containing plates as cells lacking Atf1. Therefore, residues Ser-152, Ser-172, Thr-204, Thr-216, Ser-226, and Thr-249 are essential for Atf1 function in oxidative stress survival.

Analysis of the transcriptional activity of the Atf1 phosphorylation mutants: Characterization of two subsets of stress genes

We then tested how transcription of stress genes was affected in cells expressing the phospho-Atf1 mutants. As shown by Northern blotting in Fig. 2A, cells expressing the hypophosphorylation mutant HA-Atf1.10M are not able to fully trigger the
and srx1 genes after H2O2 stress, whereas only minor defects are observed regarding activation of hsp9 and gpd1 (Fig. 2A). Therefore, the impaired activation of only a subset of genes in cells expressing the Atf1.10M mutant, including that encoding catalase, causes a severe defect in tolerance to oxidative stress; this is in agreement with our previous observation indicating that Δctt1 cells are as sensitive to peroxides as Δatf1 cells and that overexpression of just catalase is sufficient to totally suppress the sensitivity to peroxides of cells lacking Atf1 (19).

Expression of the phosho-mimicking HA-Atf1.10D mutant allows stress-dependent activation of ctt1 and srx1 to the same extent as wild-type cells; however, it constitutively induces expression of gpd1 and hsp9 (Fig. 2A).

To analyze the role of the MAP kinase Sty1 in the activation of stress genes by the HA-Atf1.10M and HA-Atf1.10D mutants, we expressed them in cells lacking Sty1. As shown in Fig. 2B, the capacity of HA-Atf1.10M to activate hsp9 and gpd1 after stress imposition was abolished in the absence of Sty1, whereas the expression of all stress genes in cells expressing HA-Atf1.10D was not altered by sty1 deletion. Concomitantly, although wild-type HA-Atf1 and the hypophosphorylation mutant HA-Atf1.10M were not able to restore wild-type tolerance to H2O2, expression of HA-Atf1.10D fully suppressed all stress defects of cells lacking Sty1 (Fig. 2C). From these results, we conclude that phosphorylation of Atf1 is sufficient for the constitutive activation of a subset of genes, such as hsp9 and...
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Figure 2. Identification of two sets of stress genes differentially regulated by Sty1-Atf1. A, stress-dependent transcriptional analysis of Atf1 phospho-mutants. YE cultures of strains 972 (WT), EP201 (Δatf1 + emp), EP203 (Δatf1 + HA-Atf1), EP203.10D (Δatf1 + HA-Atf1.10D), and EP203.10M (Δatf1 + HA-Atf1.10M) were treated with 1 mM H2O2 for 15 min or left untreated. Total RNA was analyzed by Northern blotting as described in Fig. 1D. B, expression of phospho-mimicking HA-Atf1.10D suppresses the transcription defects of cells lacking Sty1. Cultures of strains 972 (WT), EP288 (Δatf1 Δsty1 + emp), EP203 (Δatf1 Δsty1 + HA-Atf1), EP303.10D (Δatf1 Δsty1 + HA-Atf1.10D) and EP303.10M (Δatf1 Δsty1 + HA-Atf1.10M) were treated with 1 mM H2O2 for 15 min, and total RNA was analyzed as described in Fig. 1D. C, expression of HA-Atf1.10D suppresses the sensitivity to peroxides of cells lacking Sty1. The same cultures as in B were spotted on plates containing H2O2, as described in Fig. 18. D, representation of 246 genes whose transcript levels increase by at least 2-fold in wild-type cells upon treatment with 1 mM H2O2 for 15 min. Total RNA from strains 972 (WT), AV18 (Δsty1), EP303, EP303.10D, and EP303.10M (Δatf1 Δsty1 + HA-Atf1, HA-Atf1.10D, and HA-Atf1.10M, respectively) was sequenced using Illumina technology and analyzed as described under “Experimental Procedures.” Horizontal strips represent genes, and columns represent untreated or treated conditions of the indicated strains. The log2 changes in expression, relative to the untreated wild-type sample, are color-coded as shown in the bar. Genes were hierarchically clustered based on the expression patterns of cells expressing HA-Atf1.10D. The relative positions of some representative genes (srx1, ctt1, gpd1, and hsp9) are indicated on the right.

gpd1, whereas the expression of others (ctt1 and srx1) is still dependent on stress imposition. Importantly, expression of the phospho-mimicking HA-Atf1.10D (Fig. 2B) or HA-Atf1.6D (Fig S3, C and D) bypasses the requirement for a MAP kinase in the transcription process, which questions the direct participation of the kinase in Pol II initiation and/or elongation.

Because H2O2 treatment induces over 2-fold the expression of hundreds of genes in wild-type cells in a Sty1-dependent manner (2), we tested whether expression of HA-Atf1.10D in strain Δsty1 affects the whole oxidative stress-dependent gene expression program. We analyzed, by RNA sequencing, the global transcriptome of wild-type and Δsty1 cells as well as of Δsty1 expressing HA-Atf1, HA-Atf1.10D, or HA-Atf1.10M. As shown in Fig. 2D and supplemental Table S1, most of the stress genes activated after 15 min of 1 mM H2O2 in a wild-type background are not inducible in a Δsty1 background, and, in fact, their basal expression levels in this strain are significantly lower than in wild-type cells. Expression of HA-Atf1 or HA-Atf1.10M did not affect this pattern of expression of cells lacking Sty1. However, in Δsty1 cells expressing HA-Atf1.10D, more than half of the stress genes displayed up-regulation by stress more than 1.5-fold (ctt1 and srx1 are indicated by arrows in Fig. 2D), whereas half of the remaining genes displayed basal expression levels higher than 1.5-fold (similar to hsp9 and gpd1, also indicated in Fig. 2D). We conclude that HA-Atf1.10D significantly affects the gene expression pattern of strain Δsty1.

Recruitment of Atf1 to stress promoters is not dependent on its phosphorylation by Sty1

Because Atf1 displays constitutive nuclear localization, we aimed to determine whether the phosphorylation-driven event was to promote Atf1 binding to DNA. We first attempted to perform ChIP with our constitutively expressed HA-Atf1 mutants, but the amino-terminal HA tag was probably hindered from antibody recognition during ChIP experiments. We decided to add the tags at the carboxyl-terminal domain of Atf1 at the endogenous atf1 locus.

We first mutated seven central S/TP sites, rendering cells expressing Atf1.7M and Atf1.7D under the control of the endogenous atf1 promoter (supplemental Fig. S4A). Strains expressing wild-type Atf1 or Atf1.7M or Atf1.7D mutants displayed the same patterns of tolerance to peroxides and activation of stress genes as the constitutive amino-terminally tagged versions (supplemental Fig S4, B and C). We then introduced HA tags at the carboxyl-terminal coding regions of the atf1, atf1.7M, and atf1.7D chromosomal loci. Cells expressing the tagged Atf1 proteins displayed the expected tolerance to peroxides and gene expression profiles; the mutant proteins displayed the expected electrophoretic mobilities (Fig. S4, D–F).

We analyzed by ChIP the presence of wild-type Atf1-HA and phosphomutant derivatives at promoters of stress genes before and after H2O2. As shown in Fig. 3A, Atf1-HA, Atf1.7D-HA,
and Atf1.7M-HA are constitutively bound to the gpd1 and hsp9 promoters both before and after stress. Regarding the binding of wild-type and mutant Atf1-HA to the ctt1 and srx1 promoters, the TF seemed to be prebound under basal conditions relative to control primers, but 3- to 4-fold additional recruitment was detected after stress imposition (Fig. 3A). Importantly, the patterns of recruitment of Atf1, Atf1.7D, and Atf1.7M to stress promoters are very similar, indicating that the presence of Atf1 at promoters by itself is not sufficient to explain the transcriptional induction of stress genes.

**Figure 3. Recruitment of Atf1 to stress promoters is not dependent on its phosphorylation by Sty1.** A, Atf1-HA is recruited to the ctt1 and srx1 promoters after stress, whereas it is constitutively bound to the gpd1 and hsp9 promoters. MM cultures of strains CS38 (atf1-HA), CS38.7D (atf1.7D-HA), and CS38.7M (atf1.7M-HA) were treated with 1 mM H2O2 for 5 min or left untreated. ChIP experiments using anti-HA antibodies, coupled to quantification by real-time PCR, were performed using primers covering promoter regions (ctt1, srx1, gpd1, and hsp9). Primers of an intergenic region were used as a negative control. B and C, stress-dependent phosphorylation of Atf1 promotes accumulation of Pol II at stress genes. YE cultures of strains JFS5 (rpb3-HA), CS81.7D (rpb3-HA atf1.7D), and CS81.7M (rpb3-HA atf1.7M) were treated with 1 mM H2O2 for 5 min or left untreated, and CHIP assays were performed as in Fig. 3A. Importantly, the absence of Pcr1 abolishes the strong transcriptional activity of Atf1.7D-HA. YE cultures of strains 972 (WT), CS38.7D, and CS52.7D (Δpcr1 atf1.7D-HA) were treated with 1 mM H2O2 for 15 min or left untreated, and total RNA was analyzed as Fig. 3D. The recruitment of Atf1-GFP at most stress promoters, except srx1, is dependent on Pcr1. MM cultures of strains MS62 (atf1-GFP) and LS37 (Δpcr1 atf1-GFP) were treated with 1 mM H2O2 for 5 min or left untreated, and ChIP assays using anti-GFP antibodies were performed as in Fig. 3A. Data are presented as mean ± S.E. *p < 0.05 (Student’s t test).

We reported that the recruitment of Pol II at stress genes is stress-dependent, with accumulation of the polymerase subunits Rpb1 or Rpb3 at their ORFs after stress imposition (12). To confirm that active Atf1 favors recruitment of active Pol II at stress genes, we analyzed by ChIP the presence of the Pol II subunit Rpb3 at the ctt1 and gpd1 promoters, ORFs and untranslated regions in cells expressing wild-type Atf1, Atf1.7D, or Atf1.7M. As shown in Fig 3B and C, the presence of Pol II at bodies of stress genes correlates with the transcription profiles of cells expressing wild-type Atf1, Atf1.7D, or Atf1.7M (compare Fig 3B and C, with supplemental Fig. S4C). From these ChIP experiments, we conclude that the stress-dependent phosphorylation of Atf1 does not facilitate its recruitment to DNA but rather promotes, directly or indirectly, Pol II recruitment to stress genes.

**The role of other bZIP TFs, Pcr1 and Pap1, in the activation of stress genes.**

There are six TFs in *S. pombe* containing a bZIP DNA-binding motif, three of which have been clearly connected to the environmental stress response: Atf1, Pcr1, and Pap1. Pcr1 has been shown to form a heterodimer with Atf1 and to contribute to some of its functions (20, 21); however, the phenotypes and transcriptomes of strains lacking either Atf1 or Pcr1 do not fully overlap (9). As shown in Fig. 4A, Pcr1 is only dispensable in the activation of srx1 upon H2O2 stress, although its binding to stress promoters fully overlaps the binding of Atf1 (Fig. 4B). The role of Pcr1 in the activation of Sty1- and Atf1-dependent transcription of genes such as ctt1, gpd1, and hsp9 probably lies in the recognition of Atf1-binding sites at promoters, which can
only be accomplished when Atf1 is forming a heterodimer with Pcr1: expression of the Sty1-independent Atf1.7D-HA mutant cannot bypass the absence of Pcr1, as shown by the lack of transcription of stress genes (Fig. 4C). Finally, we analyzed by ChIP whether Atf1 binding to DNA is dependent on the presence of Pcr1; as shown in Fig. 4D, Atf1-GFP is not recruited to DNA in Δpcr1 cells, with the only exception of srx1. As with untagged Atf1, Δpcr1 cells expressing Atf1-GFP displayed defective activation of most stress genes except srx1 (supplemental Fig. S4G).

On the other hand, another bZIP TF, Pap1, specifically responds to H$_2$O$_2$ but not to other environmental signals, and some of its target genes overlap with those activated by Sty1-Atf1 (3); activation of Pap1 by H$_2$O$_2$ occurs through oxidation of several of its cysteine residues to disulfides and transient nuclear accumulation because of nuclear export inhibition (22, 23). We tested the contribution of this bZIP TF to the expression of the two Sty1- and Atf1-dependent subsets of genes by Northern blotting and determined that Pap1 is dispensable for the activation of gpd1 and hsp9 but required for ctt1 and srx1 (Fig. 5A). In fact, although HA-Atf1.10D is still capable of activating transcription of both subsets of genes in cells lacking Sty1, further depletion of Pap1 specifically abolishes the activation of srx1 and ctt1 but maintains constitutive expression of gpd1 and hsp9 (Fig. 5B).

Regarding the role of Pap1 and Atf1 at these genes, ChIP analysis indicates that the stress-dependent recruitment of Atf1 to ctt1 and srx1 promoters is dependent on Pap1. MM cultures of strains CS62 (Δtrr1), Pap1 is constitutively oxidized and bound to srx1 and ctt1 promoters. MM cultures of strains CS62 (Δtrr1, constitutively expressing oxidized nuclear Pap1) and CS79.C523D (pap1.C523D atf1-HA, expressing a constitutively reduced nuclear Pap1) were left untreated or treated with 1 mM H$_2$O$_2$ for 5 min. ChIP experiments were performed as in Fig. 3A. E. Pap1 binds to the ctt1 and srx1 promoters to the same extent in cells expressing or lacking Atf1. MM cultures of strains 972 (WT) and MS98 (Δatf1) were treated with 1 mM H$_2$O$_2$ for 15 min or left untreated, and ChIP experiments using anti-Pap1 antibodies were performed as in Fig. 3A.

Discussion

In S. pombe, the Sty1 pathway coordinates a wide anti-stress gene expression program in response, among others, to severe H$_2$O$_2$ stress. The role of Atf1 phosphorylation by Sty1 was uncertain, as it had been proposed to be relevant only as a protein stabilization factor. Here we dismiss this hypothesis by
showing that the protein levels of constitutively expressed HA-Atf1 do not change upon stress imposition. We demonstrate that the main role of activated Sty1 is phosphorylating Atf1 because a phospho-mimicking Atf1 mutant is able to engage the anti-stress transcriptional program in a Sty1-independent fashion. We also show that phosphorylation of the TF does not contribute to Atf1 recruitment to DNA but promotes transcription initiation.

Atf1 contains 11 putative sites of phosphorylation by Sty1. Amino acid substitutions of sites 5–10 (Ser-152, Ser-172, Thr-204, Thr-216, Ser-226, and Thr-249) are sufficient to render an inactive Atf1 because cells expressing HA-Atf1.6M are as sensitive to peroxides as strain Δatf1 (supplemental Fig. S3). We have modeled the structure of full-length Atf1 with the suite I-TASSER (25) to determine the relative position of these six phosphosites and the bZIP domain. According to this model (Fig. 6), the DNA-binding domain and five of the six phosphosites (located at a putative transactivation domain) would be separated by an intermediate domain. The intermediate domain is rich in positively charged amino acids, supporting a role in promoting Atf1 binding to DNA and buffering the gain of negative charges because of phosphorylation at the transactivation domain. An important conclusion of the predicted structure of Atf1 is that the relative position of the five phosphosites surface is very distant from the DNA-binding domain. This supports our experimentally based data, where phosphorylation of Atf1 does not have an effect on binding capacity of Atf1 to DNA but, rather, may create an interacting platform to facilitate posterior events in the transcription initiation process, such as recruitment of the SAGA complex or Pol II (12).

We have shown here that only a subset of genes, such as ctt1 and srx1, are severely affected by the absence of these phosphosites at Atf1. We are still puzzled by the fact that the hypophosphorylation mutants are able to promote transcription of the second subset of genes (gpd1 and hsp9) in a Sty1-dependent manner. We are investigating three possible scenarios: first, that Sty1 is phosphorylating a partner of Atf1, such as another bZIP TF; second, that Sty1, recruited to these promoters by HA-Atf1.10M or Atf1.7M, can also promote transcription activation; and third, that Sty1 can phosphorylate and activate Atf1 at either the five S/TP sites or at other non-canonical sites. Further work will be required to identify the target(s) of Sty1 at these stress promoters when Atf1 lacks all of these phosphorylation sites. Independent of the final outcome, we propose that Atf1 is constitutively bound to these promoters and that phosphorylation of Atf1 by Sty1 is sufficient to promote transcription initiation because the phospho-mimicking mutants (HA-Atf1.10D, HA-Atf1.6D, and Atf1.7D) can bypass the presence of Sty1 and trigger constitutive transcription of these genes.

Transcription of the Pap1-dependent set of genes, such as ctt1 and srx1, seems to follow a different pattern: a small fraction of these promoters are bound to Atf1 under basal conditions, as demonstrated by ChIP, even though this binding is significantly enhanced upon activation by stress. The stress-dependent new recruitment of Atf1 requires prior loading of Pap1, another bZIP TF that also becomes activated by H₂O₂, and it is independent of Atf1 phosphorylation status. Pap1 and Atf1 probably contribute synergistically to full activation of these essential antioxidant genes, and full transcriptional up-regulation is only accomplished when both TFs are loaded onto DNA.

The presence or absence of Sty1 does not affect the transcriptional activity of HA-Atf1.10D, and this allows us to conclude that the main role of the MAP kinase at promoters is Atf1 phosphorylation. Therefore, the putative participation of Sty1 itself in the transcription process of the S. pombe stress genes (as a transcription initiation or elongation factor, as proposed for the S. cerevisiae homologue HOG1) is dismissed by our experiments. Furthermore, Sty1 and Atf1 have been described to participate in processes other than activation of stress genes (homologous recombination, heterochromatin establishment, regulation of the ste11 and fbp1 genes, and so forth; for a review, see Ref. 26). Future experiments will show whether the phospho-mimicking Atf1 mutants can complement the absence of Sty1 in some or all of these biological functions.

**Experimental procedures**

**Yeast strains, plasmids, and growth conditions**

The origins and genotypes of strains used in this study are outlined in supplemental Table S2. We used integrative plasmid p428’ (27) and p428’ mutant derivatives to express HA-
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Atf1 and HA-Atf1 phosphomutants under the control of the constitutive sty1 promoter, most of which were generated by full-length gene synthesis (Genescript), as described previously (27). The plasmids were linearized and integrated by homologous recombination in two strain backgrounds: strain EP193 (atf1::natMX6), in which the entire atf1 open reading frame has been substituted by an antibiotic resistance cassette, and strain IV59 (atf1::natMX6 sty1::ura4), with an ura4 insertion deleting codons 35–166 of the sty1 open reading frame. The mutant plasmids were inserted at the leu1–32 locus of strains EP193 and IV59, yielding the EP203 series (∆atf1 expressing HA-Atf1 or mutant derivatives) and the EP303 series (∆atf1 Δsty1 expressing HA-Atf1 or mutant derivatives), respectively. To construct strains EP201 and EP288, used as negative controls, we integrated the p386' plasmid expressing only the sty1 promoter fused to HA at the leu1–32 locus of the EP193 and IV59 strains, respectively. To generate strains expressing Atf1.7M and Atf1.7D under the control of their own promoters, we first deleted the atf1 central region (deletion of codons 138–252 of the atf1 ORF) in the strain JA364 (h+ ura4.D18) with a cassette containing the ura4 gene, yielding strain MC117 (h+ atf1::ura4 ura4.D18). Next we replaced the mutated atf1 gene by recombination with a linear fragment of the wild-type and mutant atf1 ORF obtained by PCR amplification with specific primers using p428' and mutant derivatives as templates. Then we selected the uracil auxotrophic clones by resistance to 5-fluoro-orotic acid (Toronto Research Chemicals Inc.), yielding strain MC119.7M and MC119.7D. To tag atf1, atf1.7M, atf1.7D, rpb3, and pcr1 with HA or GFP, we transformed strains 972, MC119.7M, and MC119.7D with linear fragments containing the 3’ end of the genes fused to HA::natMX6 or GFP::kanMX6, obtained by PCR amplification using specific primers and the plasmids pFA6a-HA::natMX6 or pFA6a-GFP::kanMX6 (28), yielding strains CS38, CS38.7M, CS38.7D, LS37, JF5, CS81.7M, CS81.7D, and NG96. To delete the pap1, trr1, and pcr1 genes in the atf1::HA strains, we transformed the CS38, CS38.7M, and CS38.7D strains with linear fragments containing ORF::kanMX6, obtained by PCR amplification using ORF-specific primers and plasmid pFA6a::kanMX6 as a template, as described previously (22, 23), obtaining strain CS51 (∆pap1 atf1::HA) and its Atf1-1HA phospho-mutants (CS51.7M and CS51.7D), CS62 (∆ttr1 atf1::HA), and CS52.7D (∆pcr1 atf1.7D-HA). We mutagenized pap1 in the Atf1-1HA strain by crossing of strains IC2.C523D (pap1.C523D) and CS60 (atf1-HA) to obtain the CS79.C523D strain. Cells were grown in rich medium (YE) or synthetic minimal medium (MM) as described previously (29).

H\textsubscript{2}O\textsubscript{2} sensitivity assay

For survival on solid plates, S. pombe strains were grown, diluted and spotted on YE plates containing or not H\textsubscript{2}O\textsubscript{2} at 1 or 2 mM as described previously (9).

RNA analysis by Northern blotting

Total RNA from S. pombe YE cultures was obtained, processed, and transferred to membranes (30). Membranes were hybridized with a [α-32P]dCTP-labeled ctt1, hsp9, gpd1, srx1, or atf1 probe containing the complete ORFs. We used rRNA as a loading control.

S. pombe TCA extracts and immunoblot analysis

Modified TCA extracts were prepared as described previously (23). Tagged or untagged Atf1 was immunodetected with polyclonal anti-Atf1 (9) and polyclonal anti-GFP (31) and with house-made monoclonal anti-HA antiserum (12CA5). Anti-Sty1 polyclonal antibody (32) and monoclonal anti-tubulin (Sigma) were used as loading controls.

RNA sequencing and analysis

Total RNA from S. pombe MM cultures, treated for 15 min with 1 mM H\textsubscript{2}O\textsubscript{2} or left untreated, was obtained and processed as described previously (30). Libraries were prepared using the TruSeq Stranded mRNA Sample Prep Kit v2 (RS-122-2101/2) according to the protocol of the manufacturer. Briefly, 1 μg of total RNA was used for poly(A) mRNA selection using streptavidin-coated magnetic beads and subsequently fragmented to ~300 bp. cDNA was synthesized using reverse transcriptase (SuperScript II, 18064-014, Invitrogen) and random primers. The second strand of the cDNA incorporated dUTP in place of dTPP. Double-stranded DNA was further used for library preparation. dsDNA was subjected to A-tailing and ligation of the barcoded Truseq adapters. All purification steps were performed using AMPure XP beads. Library amplification was performed by PCR on a purified library using the primer mixture supplied in the kit. Final libraries were analyzed using an Agilent DNA 1000 chip to estimate the quantity and check size distribution and then quantified by qPCR using the Kapa library quantification kit (KK4835, Kapa Biosystems) prior to amplification with Illumina cBot. Sequencing was done using the HiSeq2000, single read, 50 nt, v3). The raw reads were inspected using FastQC v0.11.2 (Fastqc, v0.11.2, http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) for their quality and then mapped against the reference genome (S. pombe version EF2 from Ensembl corresponding to version ASM294v2 of PomBase 2015-04-21 (S. pombe, ASM294v2, http://fungiensembl.org/Chizhochecymyces_pombe/Info/Index) using TopHat version 2.0.14 (33)). Read counts were counted using htseq-count tool (version 0.6.1p1) (34) with the option -m intersection-strict and -s reverse and normalized using the rlog function of the DESeq2 package (35). The data discussed in this publication have been deposited in the NCBI Gene Expression Omnibus (36) and are accessible through GEO series accession number GSE97057. The heatmap of the rlog values was obtained by using the function pheatmap (1.0.8) from R package without reordering or clustering.

Chromatin immunoprecipitation

Cells were grown in minimal or rich medium, as indicated, and chromatin isolation and immunoprecipitation were performed as described previously (12), with the following minor changes. Cell cultures were cross-linked for 10 min instead of 20 min. After chromatin isolation as indicated (12), specific antibodies (5 μl of anti-HA antiserum (12CA5) or 1 μl of poly-
clonal anti-Pap1 or anti-GFP) were added, and incubation proceeded at room temperature while rotating for 4 h to shorten the protocol (anti-HA or anti-Pap1).

**Modeling and evaluation of the position of the six phosphosites in Atf1**

We modeled five potential conformations of the full-length sequence of Atf1 with the suite I-TASSER (25). Only two of five had a bZIP domain in the C-terminal region, and of these, we selected the structure with Thr-249 in a different domain than the rest of phosphorylatable residues (Ser and Thr). We used MODELLER (37) to model the structure of the complex with DNA formed by Pcr1 and the bZIP domain of Atf1 based on the template conformation of a cAMP-response element-binding protein–bZIP complex (38) (Protein Data Bank code 1DH3 (39)). The complete model of Atf1 and Pcr1 was obtained by substituting the bZIP domain of the model selected from I-TASSER with the model of this region obtained with MODELLER. The complex was refined with ROSETTA (40).

**Author contributions**—C. S. C., E. P., L. S. M., and M. C. conducted most of the experiments. E. P. generated most of the strains expressing Atf1 mutants and the Western and Northern blot experiments. C. S. C. performed the RNA sequencing experiments, and C. S. C. and L. S. M. performed the ChIP experiments. B. O. performed the protein modeling experiments. C. S. C., E. P., L. S. M., J. A., and E. H. designed the experiments and analyzed the results. E. H. wrote most of the paper.

**Acknowledgments**—We thank the Genomic and Bioinformatic units of the Centre for Genomic Regulation (Barcelona, Spain) for help with the performance and analysis of the RNA sequencing experiments.

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