The Yeast Snt2 Protein Coordinates the Transcriptional Response to Hydrogen Peroxide-Mediated Oxidative Stress

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The Yeast Snt2 Protein Coordinates the Transcriptional Response to Hydrogen Peroxide-Mediated Oxidative Stress

Lindsey A. Baker,* Beatrix M. Ueberheide,b Scott Dewell, Brian T. Chait, Deyou Zheng, C. David Allisa

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Regulation of gene expression is a vital part of the cellular stress response, yet the full set of proteins that orchestrate this regulation remains unknown. Snt2 is a Saccharomyces cerevisiae protein whose function has not been well characterized that was recently shown to associate with Ecm5 and the Rpd3 deacetylase. Here, we confirm that Snt2, Ecm5, and Rpd3 physically associate. We then demonstrate that cells lacking Rpd3 or Snt2 are resistant to hydrogen peroxide (H2O2)-mediated oxidative stress and use chromatin immunoprecipitation followed by high-throughput sequencing (ChIP-seq) to show that Snt2 and Ecm5 recruit Rpd3 to a small number of promoters and in response to H2O2, colocalize independently of Rpd3 to the promoters of stress response genes. By integrating ChIP-seq and expression analyses, we identify target genes that require Snt2 for proper expression after H2O2. Finally, we show that cells lacking Snt2 are also resistant to nutrient stress imparted by the TOR (target of rapamycin) pathway inhibitor rapamycin and identify a common set of genes targeted by Snt2 and Ecm5 in response to both H2O2 and rapamycin. Our results establish a function for Snt2 in regulating transcription in response to oxidative stress and suggest Snt2 may also function in multiple stress pathways.

The ability of cells to respond to stressful conditions is crucial to survival and involves both the rapid activation of stress defense proteins to begin detoxifying or protecting against the stress and gene expression changes, which bolster the stress response but take longer to manifest. In the budding yeast, Saccharomyces cerevisiae, numerous types of stress, including heat shock, oxidative stress, and nitrogen starvation, promote similar changes in a core set of genes called the "environmental stress response" (ESR) genes (1, 2). As part of the ESR, genes involved in RNA processing, translation, and ribosomal biogenesis are repressed, while genes involved in carbohydrate metabolism, protein folding, detoxification of reactive oxygen species (ROS), and maintaining redox balance are activated. These changes redirect energy away from basic metabolism toward stress detoxification and adaptation.

While the ESR is enacted similarly in response to diverse stresses, the levels and timing of gene expression changes vary depending on the type of stress, and distinct stresses promote additional changes in unique sets of genes (1). For example, while genes involved in ROS detoxification are all activated as part of the ESR, these genes are activated at higher levels when cells are exposed to oxidative stress through treatment with H2O2 (1). The stress-specific differences in the timing and extent of gene expression changes suggest that complex mechanisms regulate the ESR.

While numerous chromatin and transcriptional regulators have been linked to stress gene regulation (3–5), the full complement of proteins involved in this crucial function remains unknown. The yeast Snt2 protein was previously found to be enriched at promoter regions (6), suggesting that Snt2 may regulate transcription. In addition, Snt2 was recently reported to interact with Ecm5 and the Rpd3 deacetylase (7). Ecm5 was identified in a screen for cell wall mutants (8), but like Snt2, it contains protein domains expected to associate with chromatin (Fig. 1A). Rpd3 is a lysine deacetylase known to associate with two other well-characterized complexes, Rpd3 large [Rpd3(L)] and Rpd3 small [Rpd3(S)] (9–11). The Rpd3(L) complex has been reported to function in numerous pathways and notably has been linked to ESR regulation (12–16). However, the function of Snt2, either on its own or in association with Ecm5 and Rpd3, remains unclear.

Multiple lines of evidence point to a role for Snt2 in the oxidative stress response or possibly in the ESR more generally. First, deletion of Snt2’s reported interaction partner ECM5 results in synthetic sickness when combined with deletion of ASK10 (17), a gene originally identified as a high-copy-number enhancer of the Skn7 stress transcription factor that is involved in the oxidative stress response (18, 19). In addition, snt2Δ cells have increased phosphorylation of Slt2 (20), a mitogen-activated protein kinase (MAPK) phosphorylated in response to oxidative and other stresses (21). Finally, an integrated pathway analysis based on published stress-induced transcription changes and Snt2 genomic localization data reported a link between Snt2 and the osmotic stress response (22).

We set out to determine whether Snt2 was involved in regulating the transcriptional response to oxidative stress. We first confirmed that Snt2 physically associated with Ecm5 and Rpd3 and found that cells lacking Snt2 or Rpd3 were resistant to H2O2-mediated oxidative stress. To better understand how Snt2 and Ecm5 might function during stress, we used chromatin immuno-
were created by targeting PCR-amplified protein A (PrA), green fluorescent HygMX4 marker through homologous recombination. Tagged strains KanMX4 using standard genetic methods (23). Deletion strains were obtained from Open Biosystems. Otherwise, strains were generated by strain and are isogenic with BY4741. Where noted, deletion strains were used in this work (Table 1) were derived from the S288C background CSM from MP Biomedicals).

For rapamycin experiments, cultures were grown in synthetic defined medium supplemented with complete supplement mixture (synthetic de- tergent mixture) and used to make 5 × 10^6 cells/ml and used to make 5-fold serial dilutions. Four microliters of each dilution were spotted onto plates supplemented as indicated in the figure legends. The plates were poured no more than 20 h prior to spotting. H_2O_2 and precipitation followed by high-throughput sequencing (ChIP-seq) to map the genomic localizations of each protein before and 30 min after H_2O_2 treatment, and we identified two categories of promoters targeted by Snt2 and Ecm5, promoters very highly enriched for Snt2 and Ecm5 regardless of stress, at which both proteins were required to recruit Rpd3, and promoters at which Snt2 and Ecm5 localized independently of Rpd3 in response to H_2O_2 stress. RNA-sequencing (RNA-seq) analysis identified target genes that required Snt2 for proper expression changes in H_2O_2 stress. Including genes involved in the ESR. Finally, the snt2Δ strain was also resistant to nutrient stress imparted by the TOR (target of rapamycin) pathway inhibitor rapamycin, and rapamycin treatment enhanced Snt2 and Ecm5 localization to a subset of the H_2O_2 target promoters. Our results identify a role for Snt2 in regulating transcription in response to oxidative stress and further suggest that Snt2 may have broad roles in stress gene regulation.

MATERIALS AND METHODS

Strains, growth conditions, and growth assays. All S. cerevisiae strains used in this work (Table 1) derived from the S288C background strain and are isogenic with BY4741. Where noted, deletion strains were obtained from Open Biosystems. Otherwise, strains were generated by using standard genetic methods (23). Deletion strains were constructed by replacing the gene of interest with a PCR-amplified KanMX4 or HygMX4 marker through homologous recombination. Tagged strains were created by targeting PCR-amplified protein A (PrA), green fluorescent protein (GFP), or 13Myc (13 copies of the Myc epitope) tags to the 3’ ends of genes of interest, as previously described (23, 24). All strains were confirmed by PCR. Except where otherwise indicated, cells were grown in standard YPD medium (1% yeast extract, 2% peptone, and 2% glucose). For rapamycin experiments, cultures were grown in synthetic defined medium supplemented with complete supplement mixture (synthetic defined medium plus amino acid supplement mixture CSM [SD CSM]; CSM from MP Biomedicals).

For plate spotting assays, saturated overnight cultures of each strain were diluted in YPD (for H_2O_2 plate assays) or SD CSM (for rapamycin plate assays) and grown for 5 to 6 h to mid-log phase (optical density at 600 nm [OD_{600}] = 0.6). The strains were then diluted to 5 × 10^6 cells/ml and used to make 5-fold serial dilutions. Four microliters of each dilution was spotted onto plates supplemented as indicated in the figure legends. The plates were poured no more than 20 h prior to spotting. H_2O_2 and

![FIG 1 Snt2 associates with Ecm5 and the Rpd3 deacetylase. (A) Domain structures of Snt2, Ecm5, and Rpd3. BAH, bromo-adjacent homology; PHD, plant homeodomain finger; SANT, Sp3–Ada3–N-CoR–TFIIS (N-CoR stands for nuclear receptor corepressor 1, and TFIIS stands for transcription elongation factor IIIS); ARID, AT-rich interaction domain; JMjC, Jumonji C; HDAC, histone deacetylase. (B) Silver-stained SDS-polyacrylamide gels of eluents from Snt2-PrA and control (no tag) affinity purifications. Proteins identified by LC-MS are listed next to bands of the appropriate size. Rpd3 comigrated on the gel with IgG. Numbers to the left are molecular mass in kilodaltons. (C) Coomasie blue-stained gel analysis of Ecm5-PrA and control purifications. Proteins identified by LC-MS are listed next to the corresponding bands. (D) Lysates of unstaged, Ecm5-13Myc, Snt2-GFP, or Ecm5-13Myc Snt2-GFP strains were immunoprecipitated with anti-Myc antibody, and inputs and immunoprecipitates (IPs) were immunoblotted (IB) with anti-Myc (to detect Ecm5), anti-GFP (to detect Snt2), and anti-Rpd3, or anti-Sin3.

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* All strains isogenic to strain BY4741.
rapamycin plate assays were imaged after 2 and 3 days, respectively. Survival assays were performed as previously described (25), except that survival was determined after treatment with 0.4 mM H$_2$O$_2$ for 4 h.

Affinity purification of Snt2, EcM5, and Rpd3. *S. cerevisiae* BY4741 and cultures of strains in which Snt2 was tagged with PrA at its C terminus (Snt2-PrA), EcM5 was tagged with PrA at its C terminus (EcM5-PrA), or Rpd3 was tagged with PrA at its C terminus (Rpd3-PrA) were prepared for lysis and lysed under cryogenic conditions using a Retsch PM 100 planetary ball mill as described previously (26). Twenty grams of each lysate was resuspended in 100 ml lysis buffer (20 mM HEPES [pH 7.4], 2 mM MgCl$_2$, 300 mM NaCl, 0.1% Triton X-100, 0.1% Tween 20, 110 mM potassium acetate, 0.1 mg/ml phenylmethanesulfonyl fluoride [PMSF], 2 mg/ml pepstatin, 0.5% protease inhibitor cocktail for fungal and yeast cells [Sigma]), homogenized for 10 s with a Polytron homogenizer, incubated for 10 min with 600 units of recombinant DNase I (Roche), and clarified by centrifugation. Clarified lysates were mixed with equilibrated Dynabeads (Invitrogen) that had been conjugated with rabbit IgG as described previously (27). The rest of the purification was performed as described previously (28).

Mass spectrometric analysis of affinity purifications. For Snt2-PrA, Rpd3-PrA, and control purifications, the eluents were concentrated to dryness using a Speedvac. The eluents were resuspended in 100 mM amoniacal bicarbonate, reduced with 20 mM dithiothreitol for 4 h at 57°C, and alkylated with 45 mM iodoacetamide for 45 min in the dark at room temperature. The samples were digested with sequencing-grade modified trypsin (Promega) for 8 h at 37°C, and digestion was stopped by acidifying the solution with glacial acetic acid. The solution was pressure loaded onto self-packed precolumns (360 by 75 μm), rinsed with 0.5% acetic acid to remove salt and buffer connected to a nano-high-performance liquid chromatography (nano-HPLC) column with integrated 15-μm emitter (360 by 75 μm Picotip emitter; New Objective) packed with 6-cm of 5-μm C18 beads (YMC ODS AQ). The peptides were eluted with a linear gradient of 0 to 40% solvent B in 50 min and 40 to 100% solvent B in 70 min (solvent A is 0.1 M acetic acid, and solvent B is 70% acetonitrile in 0.1 M acetic acid) using an Agilent 1100 binary HPLC and analyzed on a Finnigan LTQ-XT mass spectrometer (Thermo Fisher) equipped with a nano-HPLC microelectrospray ionization source. The mass spectrometer was operated in a data-dependent mode where one full-scan mass spectrum was followed by 10 collision-activated dissociation (CAD) mass spectra of the 10 most abundant ions. The fragmented ions were set on an exclusion list for 40 s, and the cycle repeated throughout the data acquisition. The resulting spectra were searched against the *Saccharomyces cerevisiae* database using the searching algorithm X! Tandem (29). Proteins that have been previously reported to be contaminants of PrA affinity purifications (7, 30), that were identified in a control purification from an untagged cell lysate, or that are known to be highly abundant in yeast cells (31) were assumed to be contaminants.

For the EcM5-PrA purification, eluents were reduced and alkylated as described above, immediately separated by SDS-PAGE, and stained with Coomassie blue. Stained bands were excised, destained with 50% methanol in 100 mM ammonium bicarbonate, dehydrated, and digested overnight at room temperature in 100 mM ammonium bicarbonate with 50 ng sequencing-grade modified trypsin. Digestions were stopped by adding an aqueous solution of 5% formic acid, 0.2% trifluoroacetic acid (vol/vol), and reverse-phase resin (POROS 20 R2; Perseptive Biosystems). After light shaking at 4°C for 4 h, the resin was washed with 0.5% acetic acid, and bound peptides were eluted first with 40% acetonitrile and then with 80% acetonitrile in 0.5% acetic acid. The eluents were combined and concentrated in a Speedvac. The concentrate was pressure loaded onto a nano-HPLC column, and peptides were separated and identified as described above. The resulting spectra were searched using Sequest in the Thermo Fisher Proteome Discoverer software package (Thermo Fisher). For further validation of the Rpd3-PrA purification, 50% of the final elution was treated as described for the EcM5-PrA immunoprecipitation (IP), with focus on sequencing the bands with the molecular weights of EcM5 and Snt2, and spectra were searched using the X! Tandem search algorithm.

Immunoprecipitations and immunoblotting. For the immunoprecipitations, strains were grown to mid-log phase and collected by centrifugation. Cells were resuspended in the lysis buffer described for the affinity purifications, and lysates were prepared by glass bead disruption and clarified by centrifugation. A portion of each clarified lysate was reserved for input. For EcM5-Myc IP, the remaining lysates were incubated with anti-c-Myc beads (Sigma) for 1 h at 4°C and washed and eluted as described for the affinity purifications. For Rpd3 IP, lysates were incubated with either anti-Rpd3 antibody (sc6655; Santa Cruz) or with goat IgG (sc2028; Santa Cruz) for 1 h at 4°C, after which protein G Dynabeads (Invitrogen) were added and allowed to incubate for another hour at 4°C. Rpd3 IP were washed as described for EcM5-Myc IP. The following antibodies were used for immunoblotting: anti-MYC 9E10 (catalog no. 05-419; Millipore), anti-GFP B2 (catalog no. sc9996; Santa Cruz), anti-Rpd3 (catalog no. sc6655; Santa Cruz), anti-Snt3 (catalog no. sc17637; Santa Cruz), and anti-beta actin (catalog no. ab8224; Abcam).

Chromatin immunoprecipitation. For the H$_2$O$_2$ ChIP seq, *S. cerevisiae* BY4741 and strains in which Snt2 was tagged with 13 copies of the Myc epitope (Snt2-Myc) or EcM5 was tagged with 13 copies of the Myc epitope (EcM5-Myc) were grown to mid-log phase (OD$_{600}$ = 0.4), and an aliquot of each culture was removed and fixed for ChIP. The remaining cultures were treated with H$_2$O$_2$ (final concentration of 0.4 mM) for 30 min and then fixed similarly to the untreated samples. All samples were fixed by adding formaldehyde to a final concentration of 1% and incubating at room temperature for 20 min with rotation. Fixation was quenched by adding glycine to a final concentration of 125 mM. Cells were then pelleted, washed 4 times in cold phosphate-buffered saline (PBS), flash frozen, and stored at −80°C.

For the EcM5/Snt2-Myc rapamycin ChIP seq, cultures of each strain were grown to mid-log phase in SD CSM medium, and an aliquot was taken and fixed as described above. Each culture was then split into two cultures: one culture was treated with dimethyl sulfoxide (DMSO), while the other was treated with rapamycin dissolved in DMSO (30 nM [final concentration]). After 30 min, these cultures were fixed for ChIP as described above.

ChIP was performed essentially as described previously (32). Briefly, after lysing cells using 0.5-mm zirconia/silica beads (Biospec Products), chromatin was sonicated using a Bioruptor sonicator to 150- to 500-bp fragments. Chromatin was incubated overnight at 4°C with the Myc 9E10 monoclonal antibody (catalog no. 05-419; Millipore) and then for 2 h with protein G Magna ChIP beads (Millipore). After the DNA was washed and eluted, input and ChIP DNA samples were incubated for 15 h at 65°C to reverse cross-links and then purified using Qiagen PCR purification columns per the manufacturer’s instructions. Samples were incubated with RNase at 37°C for 2 h and then purified over Qiagen columns a second time.

Sequencing libraries of input and ChIP DNAAs were prepared using the Illumina TruSeq DNA kit per the manufacturer’s instructions except that 4% of recommended DNA adapter index concentration and 20% of the recommended PCR primer cocktail concentration were used per library. For the final amplification step, 20 and 21 cycles of PCR were used for the H$_2$O$_2$ and rapamycin libraries, respectively. Sequencing was performed on an Illumina HiSeq 2000, and data were analyzed following the Illumina pipeline.

Independent ChIP replicates were performed as described above and analyzed by quantitative PCR (qPCR) on an Applied Biosystems StepOnePlus real-time PCR system using Power SYBR green PCR master mix (Applied Biosystems) per the manufacturer’s instructions. For Rpd3-Myc ChIP experiments, cells were fixed for 45 min at room temperature in a solution consisting of 10 mM dimethyl adipimidate (DMA) and 0.25% DMSO in PBS, followed by fixation in 1% formaldehyde for 12 h. Relative enrichment was determined by dividing the percent input at the locus of interest by the percent input at the right arm of telomere 6, which had low
but detectable levels of enrichment for Snt2, Ecm5, and Rpd3, and served as a control for IP efficiency. Sequences of primers used for qPCR are available upon request.

**RNA sequencing.** BY4741, ecm5Δ, and snt2Δ cultures were treated with H2O2, as described for ChIP-seq experiments. RNA was isolated by hot acidic phenol extraction (33). Sequencing libraries were prepared from 4 μg total RNA using an Illumina TruSeq RNA kit per the manufacturer’s instructions. For the final amplification step, 15 cycles of PCR were used. An aliquot of each RNA sample was also reverse transcribed into cDNA using the SuperScript III first-strand synthesis system (Invitrogen), and RNA-seq expression values were confirmed by qPCR. Expression qPCRs were performed as described above for ChIP-qPCRs, and the levels of expression of genes of interest were normalized to the levels of ACT1 expression.

**Sequencing analysis.** ChIP-seq reads were aligned to the sacCer2 assembly of the S. cerevisiae genome using the Bowtie alignment software (34). Only unique reads that mapped to a single location with no more than 2 mismatches were kept. Regions enriched for Snt2 or Ecm5 binding (i.e., peaks) were called using the MACS (model-based analysis of ChIP-Seq) algorithm (35). Overlapping peaks were defined as peaks whose chromosomal coordinates overlapped by at least 200 bp, as determined using the Galaxy Server (36–38). For Snt2/Ecm5 correlation scatterplots, for each peak of either Snt2 or Ecm5 enrichment, we computed the Pearson’s correlation coefficients. To determine peaks where Snt2/Ecm5 enrichment increased or decreased following H2O2 treatment, a list of all peaks of shared Snt2/Ecm5 enrichment before or after treatment was compiled, and RPKM values were calculated as described above. The peaks were then separated into the peaks with Snt2 and Ecm5 RPKM values 1.5-fold higher, 1.5-fold lower, or unchanged following H2O2 treatment. Gene ontology (GO) analysis was performed using the FuncAssociate program (40). Yeast transcription start site (TSS) and open reading frame (ORF) locations were obtained by querying the Ensembl database (www.ensembl.org). The promoters were defined as the regions from 500 bp upstream of queried TSSs to the TSSs. The coordinates of peak summits (determined by the MACS algorithm) were used to uniquely assign each peak to a promoter, ORF, or neither. To determine average Snt2/Ecm5 enrichment around yeast TSSs, a custom script was used to divide regions surrounding each TSS into 50-bp windows and count the number of reads at all genes within those windows per million mapped reads. For motif analysis, 100-bp regions centered around peak summits were analyzed using the MEME suite (41). The YEASTRACT program (42) was used to search for transcription factors known to regulate Snt2/Ecm5 target genes.

RNA-seq data were aligned using the software TopHat (43), and gene expression levels and differences were calculated using the Cufflinks and Cuffdiff programs (44). All sequencing tracks were displayed using the Integrative Genomics Viewer (IGV) (45).

**Sequencing data accession number.** All ChIP-seq and RNA-seq data were submitted to the Gene Expression Omnibus (GEO) (http://www.ncbi.nlm.nih.gov/geo/) database under the accession number GSE43002.

### RESULTS

**Snt2 physically associates with Ecm5 and the Rpd3 deacetylase.** In order to better understand the function of Snt2, we first sought to confirm a previous report that Snt2 interacted with Ecm5 and Rpd3 (7). We tagged the C terminus of Snt2 with a protein A (PrA) tag (Snt2-PrA) (24) and affinity purified Snt2-PrA and associated proteins from cryogenically prepared yeast lysates (Fig. 1B). Liquid chromatography-mass spectrometry (LC-MS) analysis identified both Ecm5 and Rpd3 in the Snt2-PrA purification and not in a control purification from an untagged strain (Table 1; see Data set S1 in the supplemental material). To further confirm these associations, we repeated our affinity purifications using a strain in which Ecm5 was tagged with PrA. Consistent with the Snt2-PrA affinity purification, Snt2 and Rpd3 were both identified copurifying with Ecm5-PrA (Fig. 1C; see Data set S1).

Rpd3 is known to associate with two well-characterized protein complexes, Rpd3(L) and Rpd3(S) (9, 10). With the exception of Rpd3, components of the Rpd3(L) and (S) complexes were not identified in Snt2- or Ecm5-PrA affinity purifications (Table 2; see Data set S1 in the supplemental material), suggesting that Snt2 and Ecm5 associate with Rpd3 independently of Rpd3(L) and Rpd3(S). We next performed an affinity purification with a PrA-tagged Rpd3 strain. LC-MS analysis of this purified strain identified 10 out of 12 and 5 out of 5 of the known Rpd3-interacting proteins from the Rpd3(L) and Rpd3(S) complexes, respectively (see Data set S1). In addition, 3 Snt2 and 3 Ecm5 peptides were also identified (see Data set S1). To overcome potential suppression of signal for Snt2 and Ecm5 peptides by more abundant Rpd3-interacting proteins, a portion of the Rpd3-PrA purification was also resolved on an SDS-polyacrylamide gel, and Coomassie blue-stained gel bands corresponding in size to Snt2 and Ecm5 were excised and analyzed by LC-MS, confirming the presence of Snt2 and Ecm5 peptides in the Rpd3-PrA purification (see Data set S1).

To further confirm these results, we generated a strain in which Ecm5 and Snt2 were tagged with 13 copies of the Myc epitope or with a GFP tag, respectively. Myc-tagged Ecm5 (Ecm5-Myc) was immunoprecipitated using a Myc antibody, and immunoblot analysis confirmed that GFP-tagged Snt2 (Snt2-GFP) and Rpd3 both coprecipitated with Ecm5-Myc (Fig. 1D). In contrast, Sin3, a component of the Rpd3(L) and Rpd3(S) complexes, did not co-precipitate with Ecm5-Myc (Fig. 1D). Taken together, our results confirm that Snt2, Ecm5, and Rpd3 physically associate in a form separable from Rpd3(L) and Rpd3(S).

**Cells lacking Snt2 or Rpd3 are resistant to H2O2-induced oxidative stress.** We next sought to determine the function of Snt2, either alone or in association with Ecm5 and Rpd3. A recent genetic screen found that deletion of ECM5 resulted in synthetic sickness when combined with deletion of the ASK10 gene (17), which encodes a protein involved in the oxidative stress response (18). Furthermore, cells lacking Snt2 have higher levels of phosphorylation of the stress MAPK Stt2 (20), and a bioinformatics

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d Proteins also identified in control purification or known to be contaminants of PrA purifications (7, 30) were omitted.

b Phosphorylated peptide confirmed by tandem MS (MS-MS).

c While not previously reported as contaminants, these proteins are highly abundant (31) and are therefore likely contaminants.
d Log of E score generated by X!Tandem algorithm (29).
study suggested that Snt2 might function to activate genes in response to stress (22). Rpd3 is known to regulate the response to numerous stresses, including oxidative stress, often as a part of the Rpd3(L) complex (12–15). On the basis of these data, we hypothesized that Snt2, Ecm5, and Rpd3 might function as an additional regulator of the oxidative stress response.

To test this hypothesis, we spotted serial dilutions of wild-type, snt2Δ, ecm5Δ, and rpd3Δ strains on untreated or H$_2$O$_2$-treated plates. While all the strains grew similarly on untreated plates (with the rpd3Δ strain having a slight growth defect), there was marked contrast in the growth of these strains on plates containing H$_2$O$_2$ (Fig. 2A). The wild-type and ecm5Δ strains showed strong H$_2$O$_2$ sensitivity. Consistent with the known role for Rpd3 in regulating the yeast oxidative stress response as part of the Rpd3(L) complex, the rpd3Δ strain was resistant to H$_2$O$_2$. Strikingly, the snt2Δ strain was even more resistant to H$_2$O$_2$ than the rpd3Δ strain, showing similar levels of resistance to a strain lacking Gpr1, a glucose sensor whose deletion was reported to result in H$_2$O$_2$ resistance of double deletion strains for these factors. Surprisingly, the strain having a slight growth defect, there was also more Rpd3 precipitating with Ecm5-Myc before and after H$_2$O2 treatment (Fig. 3A). Separately derived snt2Δ and rpd3Δ strains constructed on the BY4742 background also showed H$_2$O$_2$ resistance in this assay (data not shown).

We next determined whether snt2Δ and rpd3Δ strains also showed H$_2$O$_2$ resistance using a liquid survival CFU assay (25). When wild-type or ecm5Δ cells were treated with H$_2$O$_2$ at a final concentration of 0.4 mM for 4 h, approximately 20% of cells survived (Fig. 2B). In contrast, the rpd3Δ strain had enhanced survival, with an average survival rate of 40.7%. Moreover, greater than 50% of snt2Δ cells survived H$_2$O$_2$ treatment, a statistically significant difference from the wild-type levels, confirming that cells lacking Snt2 are resistant to oxidative stress.

The differing H$_2$O$_2$ sensitivities of the snt2Δ, rpd3Δ, and ecm5Δ mutants suggested that despite their physical association, these proteins might have distinct functions in the oxidative stress pathway. In order to better understand how these proteins function to regulate stress tolerance, we next determined the H$_2$O$_2$ resistance of double deletion strains for these factors. Surprisingly, deletion of ECM5 reversed the H$_2$O$_2$ resistance seen in the snt2Δ strain (Fig. 2C), suggesting that Ecm5 has a function in the oxidative stress response that is opposed to the function of Snt2. Although the snt2Δ rpd3Δ double knockout possessed a modest growth defect similar to that of the rpd3Δ strain (as can be seen by the smaller size of the snt2Δ rpd3Δ colonies in Fig. 2C), the double knockout was more H$_2$O$_2$ resistant than the rpd3Δ strain alone, showing similar levels of resistance to the snt2Δ strain. This result suggests that deletions of RPD3 and SNT2 promote stress resistance through different pathways, with the stress resistance of the rpd3Δ strain possibly relating to the known role of the Rpd3(L) complex in regulating the oxidative stress response. In agreement, strains lacking the Rpd3(L) complex members Sin3, Sap30, and Cts6 were also H$_2$O$_2$ resistant (Fig. 2D). Taken together, these results suggest that even though Snt2, Ecm5, and Rpd3 physically associate in cells, Snt2 performs a function in the oxidative stress response distinct from that of Ecm5 or Rpd3.

Similar levels of Snt2, Ecm5, and Rpd3 associate with one another before and after H$_2$O$_2$ treatment. We next determined whether oxidative stress affected the associations between Snt2, Ecm5, and Rpd3. Immunoblot analysis of Ecm5-Myc, Snt2-GFP, and Rpd3 protein levels in Myc immunoprecipitations from ECM5-MYC SNT2-GFP or SNT2-GFP lysates did not reveal significant differences in the levels of Snt2-GFP and Rpd3 associating with Ecm5-Myc before and after H$_2$O$_2$ treatment (Fig. 3A). Similarly, when Rpd3 was immunoprecipitated from ECM5-MYC SNT2-GFP lysates, there were not differences in the levels of Snt2-GFP and Rpd3 associating with Ecm5-Myc before and after H$_2$O$_2$ treatment (Fig. 3A). Similarly, when Rpd3 was immunoprecipitated from ECM5-MYC SNT2-GFP lysates, there were not differences in the levels of Snt2-GFP and Ecm5-Myc coprecipitating with Rpd3 before and after H$_2$O$_2$ stress (Fig. 3B [note that while there was slightly more Ecm5-Myc and Snt2-GFP detected in the immunoprecipitation from the H$_2$O$_2$-treated cells, there was also more Rpd3 precipitated]). Thus, H$_2$O$_2$ stress does not result in changes in the levels of Snt2, Ecm5, and Rpd3 physically associating with each other.

In response to H$_2$O$_2$ stress, Snt2 and Ecm5 colocalize to gene promoters. To better understand how Snt2 and Ecm5 function in the oxidative stress response, we used ChIP-seq to map the genomic localizations of Myc-tagged Snt2 or Ecm5 both before and after H$_2$O$_2$ treatment. We chose to focus on Snt2 and Ecm5 to avoid having to distinguish Rpd3 localization associated with Snt2 and Ecm5 from Rpd3 localization via the Rpd3(L) and
Rpd3(S) complexes. Consistent with their physical association, the Snt2 and Ecm5 ChIP profiles were strikingly similar (representative region shown in Fig. 4A). Both before and after H2O2 treatment, the majority of peaks of Snt2 enrichment overlapped with Ecm5 peaks by at least 200 bp (Fig. 4B). Scatterplots confirmed correlations between Snt2 and Ecm5 enrichment levels (Fig. 4C, Pearson’s correlations of 0.895 and 0.917 for the 0-min and 30-min data sets, respectively).

We next analyzed the genomic distributions of Snt2 and Ecm5 peaks for enrichment at particular genetic features (Fig. 4D). Before treatment, 43% of peaks fell in promoter regions, while 23% fell within open reading frame (ORF) regions. Reads from an untagged control strain were also enriched in ORF peak regions (data not shown), suggesting that the ORF peaks may be an artifact reflecting the higher accessibility of transcribed chromatin rather than real sites of Snt2/Ecm5 enrichment, an effect that has been previously reported for ChIP experiments involving other proteins (48). We therefore chose to focus on peaks localized to promoter regions for further study. Strikingly, 30 min after H2O2 treatment, Snt2 and Ecm5 underwent dramatic localization changes, with many regions showing new or enhanced localization of Snt2 and Ecm5 (Fig. 4A). After H2O2 treatment, 74% of peaks were in promoters (Fig. 4D). Alignments of Snt2 and Ecm5 ChIP-seq reads around the transcription start sites (TSSs) of all genes revealed Snt2 and Ecm5 enrichment approximately 250 bp upstream of the TSS in H2O2-treated cells (Fig. 4E, dark purple lines). In addition, there were 817 peaks of Snt2 and Ecm5 enrichment identified after treatment, compared to only 315 peaks present before (Fig. 4B). For both Snt2 and Ecm5, the majority of peaks called in untreated samples were also present after H2O2 treatment (Fig. 4F).

In order to compare between different ChIP-seq data sets, we generated a list of all peaks of both Snt2 and Ecm5, identified before or after stress, and calculated Snt2 or Ecm5 enrichment at each region before and after stress (see Materials and Methods). When all peaks were considered together, enrichment of Snt2 and Ecm5 was significantly higher after treatment compared to before treatment, consistent with the many new peaks identified in treated cells (Fig. 4G). Similarly, scatterplots showing Snt2 or Ecm5 enrichment at all peak regions before treatment compared to enrichment after treatment show that the majority of peaks had enrichment scores at least 1.5-fold higher after H2O2 treatment compared to enrichment before treatment (Fig. 4H [note that purple points outnumber gray and orange points]).

To look for motifs in Snt2/Ecm5 target sites, we used the de novo motif finder Meme (41). The 100 bp centered around each peak’s summit (determined by the MACS algorithm) was used for this analysis. While no motif was significantly enriched when all peaks were analyzed together, when we used the 20 peaks with the highest Snt2/Ecm5 enrichment levels for this analysis, a motif containing (A/G)/(C/T)/GGCGCTA/(C/T)/CA was strongly enriched (Fig. 4I, E value of $1.3 \times 10^{-10}$), in agreement with an Snt2-binding motif identified in a previous Snt2 chromatin immunoprecipitation with microarray technology (ChIP-chip) study, (C/T)/GGCGCTA/(C/T)/CA (6). We next analyzed the peaks where Snt2 and Ecm5 levels were at least 1.5-fold higher after H2O2 treatment (purple points in Fig. 4H). A second motif, CCG(T/G)GA, was identified among these H2O2-enriched Snt2/Ecm5 peaks (Fig. 4I, right, E value of $1 \times 10^{-3}$). This motif is similar to the previously published motifs of the Pdr1 and Pdr3 transcription factors (CCGCGGA) (49), which regulate cellular transport genes (50). Collectively, these results show that H2O2 treatment stimulates Snt2 and Ecm5 to colocalize to promoters and identify Pdr1 and Pdr3 as potential factors that may cooperate with Snt2 and Ecm5.

After H2O2 treatment, Snt2 and Ecm5 localize to promoters of genes involved in oxidative stress and cellular metabolism. We next sought to determine which categories of genes were targeted by Snt2 and Ecm5 as a result of oxidative stress. As described above, we identified peaks in which Snt2/Ecm5 enrichment increased, decreased, or did not change after H2O2 treatment (purple, orange, or gray points, respectively, in Fig. 4H), and analyzed each set of target genes using gene ontology (GO) analysis (40). Genes whose promoters had decreased levels of Snt2 and Ecm5 after H2O2 stress were involved in protein metabolism (ribosome and translation) (Fig. 4J). Genes whose promoters had similar levels of Snt2 and Ecm5 before and after treatment also functioned in protein metabolism, as well as in carbohydrate metabolism (gluconeogenesis, glycolysis, and transposition [Fig. 4J]). Thus, in unstressed cells, Snt2 and Ecm5 target sugar and protein metabolism genes.

Genes where Snt2 and Ecm5 levels increased after H2O2 treatment had functions in amino acid, steroid, and alcohol metabolism (Fig. 4I). For example, after H2O2 treatment, Snt2 and Ecm5 were enriched at the promoters of the BAT2 amino acid metabolism gene and the sterol metabolism genes ERG3 and ERG6, which help synthesize the cell wall component ergosterol (Fig. 5A). While the cell wall category was not identified by GO analysis, we found additional cell wall genes targeted by Snt2 and Ecm5 in response to H2O2 treatment, including the cell wall glucanase SCW10 and the glucan synthase FKS3 (Fig. 5A). Ecm5 was originally identified in a screen for mutants with cell wall defects (8), and these results suggest a possible connection between Ecm5 function and cell wall maintenance. In response to stress, Snt2 and Ecm5 were also enriched at cellular transport genes (Fig. 4I) including the FUI1 uridine permease (Fig. 5A), consistent with the enrichment of a motif similar to that of the Pdr1 and Pdr3 transport gene regulators among Snt2/Ecm5 target sites. Notably,
FIG 4 Snt2 and Ecm5 are highly colocalized and associate with additional promoters after H2O2 stress. (A) ChIP-seq tracks showing Snt2 and Ecm5 ChIP enrichment in a representative region before and after H2O2 treatment. The coverage values for each track are scaled by 1,000,000/number of reads, and the scale of the y axes for all tracks is shown in brackets. The locations of genes and peaks and the chromosomal coordinates are shown under the tracks. chrII, chromosome II. (B) Venn diagrams show that the majority of Snt2 and Ecm5 ChIP peaks overlap. (C) Correlations between Snt2 and Ecm5 enrichment before and after H2O2 treatment (Pearson’s correlation coefficients are indicated in the bottom right-hand corners of the graphs). (D) Genomic distributions of shared Snt2/Ecm5 peaks before and after H2O2 treatment (Pearson’s correlation coefficients are indicated in the bottom right-hand corners of the graphs). (E) Overlaps of Snt2 or Ecm5 peaks before and after H2O2 treatment. (G) Box-and-whisker plots showing the distributions of the enrichments at all peak regions before and after H2O2 treatment in the treated and untreated data sets. The bottoms, middles, and tops of the boxes depict the 25th, 50th, and 75th percentiles, respectively, and the top and bottom whiskers depict the 10th and 90th percentiles, respectively, of Snt2 or Ecm5 genes, scaled by 1,000,000/total reads. (F) Overlaps of Snt2 or Ecm5 peaks before and after H2O2 stress. (E) Average number of Snt2 or Ecm5 ChIP-seq reads per 50-bp window around transcription start sites (TSSs) for all yeast transport genes are all categories regulated as part of the environmental stress response (ESR) (1), showing that in response to stress, Snt2 and Ecm5 target ESR genes. Consistent with genetic evidence linking Snt2 to oxidative stress regulation, genes involved in redox reactions and the oxidative stress response also had increased levels of Snt2 and Ecm5 after treatment (Fig. 4J). These genes included CYC1, which encodes the cytochrome c component of the mitochondrial electron transport chain, and SUE1, which encodes a protein that degrades ergosterol biosynthesis, amino acid metabolism, and cellular transport genes are all categories regulated as part of the environmental stress response (ESR) (1), showing that in response to stress, Snt2 and Ecm5 target ESR genes. Consistent with genetic evidence linking Snt2 to oxidative stress response, oxidative stress response; a.a. metabolism, amino acid metabolism.
unstable cytochrome c (Fig. 5A). The incomplete reduction of oxygen in the electron transport chain is one of the main endogenous sources of ROS in cells (51), and the association of Snt2 and Ecm5 with promoters of genes involved in cytochrome c biology further links these proteins to oxidative stress. In addition, after H2O2 treatment, Snt2 and Ecm5 were enriched in the promoters of the genes encoding the oxidative stress transcription factors Yap1 and Cin5 (also known as Yap4) which are activated by oxidative stress (52,53), the catalase Ctt1, and the superoxide dismutase Sod1, which detoxify ROS (54), as well as the membrane proteins Hsp30 and Hsp12, which are induced in response to oxidative stress (55,56) (Fig. 5A and data not shown). Increased Snt2 and Ecm5 localization to many of these promoters in response to H2O2 treatment was confirmed by ChIP-qPCR (Fig. 5B and data not shown).

A small number of promoters were very highly enriched for Snt2 and Ecm5. ChIP-seq analysis identified a small set of promoters very highly enriched for Snt2 and Ecm5 both before and after H2O2 treatment: while most peaks of Snt2 and Ecm5 enrichment had peak heights between 70 and 150 (representing the number of reads in the peak summit scaled by 1,000,000/total number of reads in that ChIP-seq experiment), we identified 10 “superenriched” Snt2 and Ecm5 peaks in the promoters of 14 genes, with peak heights over 500 in both untreated and treated cells (Table 3). For example, Snt2/Ecm5 peaks at the promoters of CYC3/CDC19, BNR1/POT1, and MSN1 were more than 10-fold more enriched than the peaks described in the previous section (compare the values within brackets in Fig. 6A with those in Fig. 5A). Independent ChIP-qPCR experiments recapitulated the high levels of Snt2 and Ecm5 enrichment found at these promoters (Fig. 6B), showing that these high enrichment levels are not merely sequencing artifacts.

While the only significant GO association found for the 14 superenriched target genes was polysaccharide catabolism (P = 0.0002), further molecular analysis suggested that the promoters that were superenriched for Snt2 and Ecm5 were enriched for motifs for several transcription factors known to be involved in metabolism (Fig. 6C). Further, the ChIP-seq analysis of Snt2 and Ecm5 enrichment was able to predict both the peak locations and the enrichment levels. This suggests that the ChIP-seq analysis may be able to predict which promoters are likely to be enriched for Snt2 and Ecm5 in response to oxidative stress in general.

FIG 5 After H2O2 treatment, Snt2 and Ecm5 localize to stress and metabolism genes. (A) Examples of ESR genes whose promoters are enriched for Snt2 and Ecm5 after H2O2 treatment. For bidirectional promoters, the gene associated with the category above the panel is shown in a larger font. (B) ChIP-seq results were confirmed by ChIP-qPCR. Relative enrichment was determined by normalizing percent inputs at the target locus to percent inputs at a control region on the right arm of telomere 6. The means ± SEMs of 3 biological replicates are shown. Samples in which the ChIP enrichment after treatment differs significantly from enrichment before treatment are indicated by asterisks as follows: *, P < 0.05; **, P < 0.01. Enrichment in the ACT1 ORF is shown as a negative control.
likely due to the small number of genes, almost all of the genes had functions connected to the ESR (Table 3), including genes involved in carbohydrate metabolism (GDB1, CDC19, and PGU1), fatty acid metabolism (POT1), ribosome biogenesis (IPI3), translation (STM1), amino acid transport (CUP9), redox reactions (YNL181W and CYC3), and general stress response (SSA3, CUR1, and MSN1). The high levels of Snt2 and Ecm5 at superenriched promoters may reflect high occupancy of these regions in all cells, or alternatively, a high proportion of cells in the culture that have Snt2 and Ecm5 occupied at these sites.

**Sn2 and Ecm5 are required for Rpd3 recruitment to superenriched promoters but not for recruitment of the Rpd3(L) complex member Sds3.** Because Rpd3 physically associated with Sn2 and Ecm5, we next determined whether Rpd3 also associated at Sn2/Ecm5 target sites, and if so, whether it did so in an Sn2- or Ecm5-dependent manner. Deletion of ECM5 or SNT2 from a Myc-tagged Rpd3 strain did not alter global levels of Rpd3-Myc before or after H2O2 treatment (Fig. 6C). By ChIP-qPCR, Myc-tagged Rpd3 was enriched at all Sn2/Ecm5 target promoters assayed, both before (data not shown) and after (Fig. 6D) treatment. Surprisingly, deletion of SNT2 or ECM5 abrogated Rpd3-Myc enrichment at Sn2/Ecm5 superenriched promoters but not at promoters where Sn2 and Ecm5 were enriched after H2O2 treatment (Fig. 6D). Thus, there are at least two classes of Sn2/Ecm5 targets, those at which Sn2 and Ecm5 localize at high levels regardless of stress and recruit Rpd3, and those at which Sn2 and Ecm5 associate in response to H2O2 treatment, independently of Rpd3.

Because Rpd3 recruitment can also occur through association with the Rpd3(L) complex and Rpd3(L) function has been linked to the yeast stress response (11–16), we sought additional confirmation that the Sn2- and Ecm5-dependent association of Rpd3 with superenriched promoters occurred independently of the

<table>
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<td>SSA3</td>
<td>HSP70 family chaperone involved in the stress response</td>
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<tr>
<td>3</td>
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<td>Thiolase involved in beta-oxidation of fatty acids</td>
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<td>Secreted polygalacturonase used to hydrolyze plant pectins</td>
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<td>STM1</td>
<td>Ribosome-binding protein required for translation under nutrient stress</td>
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<tr>
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<td>Glycogen debranching enzyme involved in glycogen degradation</td>
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**FIG 6** Sn2 and Ecm5 are required for Rpd3 recruitment to superenriched promoters but not for recruitment of the Rpd3(L) complex member Sds3. (A) Promoters superenriched for Sn2 and Ecm5 (note the ranges in brackets for ChIP-seq tracks) before and after H2O2 treatment. (B) ChIP-qPCR confirmation of high levels of Sn2 and Ecm5 at superenriched promoters shown in panel A. Data were normalized as described in the legend to Fig. 5B. (C) Control immunoblots showing that deletion of SNT2 or ECM5 did not affect Rpd3-Myc levels and that deletion of SNT2 did not affect Sds3-Myc levels before or after H2O2 treatment. Actin blots serve as a loading controls. (D) Rpd3-Myc ChIP-qPCR performed on untagged, RPD3-MYC, RPD3-MYC snt2Δ, or RPD3-MYC ecm5Δ strains that were treated with H2O2. Data were normalized as described in the legend to Fig. 5B. ACT1 ORF is the negative control. (E) Sds3-Myc ChIP-qPCR performed on untagged, SDS3-MYC, or SDS3-MYC snt2Δ strains that were treated with H2O2. Values are means ± standard deviations of 3 independent measurements and are representative of 2 experiments.
Rpd3(L) complex. We tagged the Rpd3(L) subunit Sds3 with the Myc tag (Sds3-Myc) and used ChIP-qPCR to analyze Sds3-Myc association with Snt2/Ecm5 target promoters in the presence or absence of Snt2. Deletion of SNT2 did not affect Sds3-Myc protein levels before or after H$_2$O$_2$ treatment (Fig. 6C). Sds3-Myc was not enriched above background levels at the MSN1 or IP13/YNL181W Snt2/Ecm5 superenriched promoters (Fig. 6E), suggesting that at these targets, Snt2, Ecm5, and Rpd3 function independently from the Rpd3(L) complex. Sds3-Myc was modestly enriched at the superenriched CYC3/CDC19 promoter. However, Sds3-Myc enrichment of these promoters and the EXG1 promoter, an H$_2$O$_2$-enriched Snt2/Ecm5 target, was not decreased when SNT2 was deleted, showing that Snt2 and Ecm5 do not recruit Rpd3(L) to target promoters (Fig. 6E).

**Snt2, but not Ecm5, is required for proper expression of ChIP target genes.** Having shown that Snt2 and Ecm5 localize to ESR genes after H$_2$O$_2$ treatment, we next sought to determine whether Snt2 or Ecm5 regulated target gene expression. We performed RNA-sequencing (RNA-seq) analysis of wild-type, snt2Δ, or ecm5Δ strains before or 30 min after H$_2$O$_2$ treatment. The RNA-seq analysis was performed on three replicate samples, and expression levels of many Snt2 and Ecm5 target genes were confirmed by qPCR analysis (data not shown). In wild-type cells, 3,127 genes significantly changed expression in response to H$_2$O$_2$ treatment (significance determined by the CuffDiff program). A previous microarray study reported 1,294 genes up- or downregulated at least 2-fold in wild-type cells 30 min after treatment with 0.32 mM H$_2$O$_2$ (1). The larger number of genes identified in our study function in acetyl coenzyme A (acetyl-CoA) metabolism, NAD metabolism, oxidation-reduction processes, and the tricarboxylic acid cycle enriched among genes whose expression changed in the snt2Δ strain (P values of 8.8 × 10$^{-7}$, 5.5 × 10$^{-5}$, 1.3 × 10$^{-6}$, and 5.8 × 10$^{-6}$, respectively). After H$_2$O$_2$ treatment, genes upregulated in the snt2Δ strain functioned in protein synthesis, rRNA processing, and hexose transport, while genes downregulated in the snt2Δ strain functioned in translation, amino acid metabolism, ROS metabolism, and response to ROS (Fig. 7C), further linking Snt2 to oxidative stress and metabolism regulation.

By integrating our RNA-seq and ChIP-seq analyses, we identified 403 of the 1,205 H$_2$O$_2$-enriched Snt2/Ecm5 target genes that were up- or downregulated at least 1.5-fold in the snt2Δ strain after treatment. Of the 403 genes, 309 genes also exhibited changes in expression in wild-type cells in response to H$_2$O$_2$ treatment (Fig. 7D), demonstrating considerable overlap between target genes directly regulated by Snt2 and target genes regulated through the oxidative stress response. Notably, many of the Snt2 and Ecm5 targets involved in the oxidative stress response or in cytochrome c regulation described above had altered expression levels in the snt2Δ strain after treatment (Fig. 7E). Taken together, these results show that H$_2$O$_2$ treatment prompts Snt2 to target and regulate stress response genes.

When the 309 target genes regulated by Snt2 were clustered on their expression profiles, 4 clusters could be distinguished, encompassing genes whose expression increased (clusters 1 and 2) or decreased (clusters 3 and 4) in wild-type cells after H$_2$O$_2$ treatment and genes whose expression levels in the snt2Δ strain after treatment were higher (clusters 1 and 3) or lower (clusters 2 and 4) than wild-type expression levels (Fig. 7F). While a regulatory trend was not readily observed when all 309 genes were considered together (Fig. 7G), distinct patterns did emerge within specific categories of genes. For instance, all but one of the 24 translation genes in this set were downregulated in the snt2Δ strain after treatment. In contrast, 38 of the 48 plasma membrane genes in this set were upregulated in the absence of Snt2, as were all 20 of the lipid metabolism genes, including the ergosterol metabolism genes.

Our expression results suggest that Snt2 functions to both promote and repress gene expression, depending on the target. A mechanism through which this might occur may be through Snt2-mediated stabilization of unique sets of transcriptional regulators at different genes. As mentioned above, a subset of H$_2$O$_2$-enriched Snt2/Ecm5 target promoters contained a motif similar to the motif of the Pdr1 and Pdr3 transcriptional regulators. To search for other transcription factors that might cooperate with Snt2, we used the YEASTRACT program (42) to search for factors with binding sites in the 309 Snt2-regulated target genes that exhibited changes in expression in wild-type cells in response to stress (Fig. 7H) and found that 205 of the 309 genes are targets of the Ste12 transcription factor, which is activated downstream of the Fus3 mating and Kss1 filamentous growth MAPK pathways and is also activated by nutrient stress (57). In addition, 138 of the 309 genes were targets of Rap1, which functions as both a transcriptional activator and repressor depending on the target (58), and regulates ribosomal biogenesis genes (59). Thus, Ste12 and Rap1 may be two additional factors that potentially cooperate with Snt2 to regulate gene expression.

**Treatment with the TOR pathway inhibitor rapamycin promotes Snt2 and Ecm5 localization to many of the same genes as H$_2$O$_2$ treatment.** The genes regulated by Snt2 and targeted by Snt2 and Ecm5 after H$_2$O$_2$ treatment had numerous functions linked to stress response, including genes involved in metabolism and nu-
trient transport. We therefore hypothesized that rather than functioning solely in the oxidative stress response, Snt2 might function more broadly to regulate stress and metabolism. To test this, we interrogated the growth of the snt2Δ/H9004 strain in the presence of rapamycin, a TOR pathway inhibitor that mimics nitrogen starvation (60). In dilution spotting assays, snt2Δ/H9004 cells were resistant to rapamycin (Fig. 8A) (the less nutrient-rich SD CSM medium was chosen for these assays to ensure that the high levels of nutrients in YPD medium did not compensate for a nutrient stress phenotype).

We used ChIP-seq to map Snt2 and Ecm5 localization in cells treated for 30 min with rapamycin or with DMSO as a control. Similar to H2O2 treatment, rapamycin treatment prompted increased Snt2 and Ecm5 association at multiple loci (marked with small black arrows in Fig. 8B). Of the 155 Snt2/Ecm5 peaks where Snt2 and Ecm5 levels increased at least 1.5-fold in response to rapamycin treatment, 116 were also regions where Snt2/Ecm5 levels increased after H2O2 treatment (Fig. 8C). For example, both H2O2 and rapamycin treatment prompted Snt2 and Ecm5 enrichment at the promoters of the FUI1 uridine permease and BAT2 amino acid aminotransferase genes (Fig. 8D). These results suggest that rather than Snt2 being a unique regulator of oxidative stress, Snt2 regulates stress and metabolism more broadly.

**DISCUSSION**

All cells must balance allocation of energy to growth with allocation to stress defense mechanisms, and when cells are exposed to stress, they must rapidly shift investment to the latter. In yeast, this shift involves regulation of a common set of genes called the ESR genes (1,2). However, the mechanisms through which these genes are regulated are not fully understood. Here, we have shown that Snt2 physically associates with Ecm5 and the Rpd3 deacetylase and have integrated phenotypic, ChIP-seq, and RNA-seq analyses to uncover an Rpd3-independent role for Snt2 in regulating genes in response to oxidative stress. While a previous study mapped Snt2 localization using ChIP-chip (6), as far as we know, ours is the first study to map Snt2 using high-throughput sequencing, and more importantly, the first to examine the localization changes of either Snt2 or Ecm5 in response to stress.

Consistent with a previous report (7), our LC-MS analysis of affinity-purified PrA-tagged Snt2, Ecm5, and Rpd3 cell lysates confirmed associations between these three proteins (Fig. 1 and...
and Rpd3 comprise a third Rpd3 complex. We identified a second set of promoters where levels of Snt2 and Ecm5 increased after H$_2$O$_2$ stress (Fig. 4H and 5). While Rpd3 was enriched at these promoters by ChIP-qPCR, deletion of SNT2 or ECM5 did not affect Rpd3 association (Fig. 6D), suggesting that other proteins recruit Rpd3 to these targets. The Rpd3(L) complex regulates genes in response to diverse stresses (13–16), including oxidative stress (Fig. 2A and B) and had few gene expression changes (Fig. 7A), but deletion of the SNT2 or ECM5 gene from the /H9004 strain reversed the H$_2$O$_2$ resistance of this strain (Fig. 2C), suggesting that these proteins exert such opposing effects remain to be elucidated, but they may involve the cooperation of unique factors at different promoters. A subset of the H$_2$O$_2$-enriched promoters contained a motif similar to the known Pdr1/Pdr3 motif (Fig. 4I), and many of the Snt2/Ecm5 target genes regulated by Snt2 are also known to be targeted by Ste12 and Rap1 (Fig. 7H), suggesting that Pdr3, Pdr1, Ste12, and Rap1 may cooperate with Snt2 to regulate different gene sets.

Surprisingly, despite the physical association and genomic colocalization of Snt2 and Ecm5, our genetic data uncovered independent functions for these proteins in the response to H$_2$O$_2$ stress. Unlike the snt2Δ strain, the ecmlΔ strain was not resistant to oxidative stress (Fig. 2A and B) and had few gene expression changes (Fig. 7A), but deletion of the ECM5 gene from the snt2Δ strain reversed the H$_2$O$_2$ resistance of this strain (Fig. 2C), suggesting that Ecm5 has a function that is opposed to that of Snt2.

Table 2: see Data set S1 in the supplemental material). While Rpd3 is known to associate with two other complexes (9, 10), we did not identify subunits of either complex associating with Snt2 or Ecm5 (Fig. 1D and Table 2; see Data set S1), suggesting that Snt2, Ecm5, and Rpd3 comprise a third Rpd3 complex.

Strikingly, both the snt2Δ and rpd3Δ strains were resistant to H$_2$O$_2$-mediated oxidative stress (Fig. 2A and B), leading us to initially hypothesize that Snt2, Ecm5, and Rpd3 function together in the oxidative stress response. We used ChIP-seq to identify regions where Snt2 and Ecm5 were localized before and 30 min after H$_2$O$_2$ treatment (Fig. 4) and surprisingly, identified at least two distinct types of Snt2/Ecm5 targets: a small number of promoter regions at which Snt2 and Ecm5 are required for Rpd3 localization (Fig. 6) and a much larger set of promoters where Snt2 and Ecm5 localize independently of Rpd3 in response to H$_2$O$_2$ treatment (Fig. 5).

In the first set of promoters, high levels of Snt2 and Ecm5 were detected in both untreated and H$_2$O$_2$-treated cells (Fig. 6A and B and Table 3). Rpd3 was enriched at these promoters, and Snt2 and Ecm5 were required for Rpd3 enrichment (Fig. 6D and data not shown). Consistent with our inability to detect Rpd3(L) subunits associating with Snt2 and Ecm5, deletion of SNT2 did not affect the localization of the Rpd3(L) complex member Sds3 (Fig. 6E), further supporting the idea that Snt2, Ecm5, and Rpd3 function as a distinct complex. While many of the superenriched promoters had functions linked to stress response, Snt2, Ecm5, and Rpd3 associated with these regions before and after H$_2$O$_2$ treatment. Furthermore, we did not detect altered levels of Snt2-GFP, Ecm5-Myc, and Rpd3 associating with one another after H$_2$O$_2$ treatment (Fig. 3), suggesting that Snt2, Ecm5, and Rpd3 may have a stress-independent function at these promoters. Surprisingly, while deletion of SNT2 or ECM5 decreased Rpd3 levels at these promoters, we did not detect changes in histone H3 acetyl levels in these regions by ChIP-qPCR (data not shown). Rpd3 may be recruited to these regions to deacetylate other histone substrates or nonhistone proteins. Alternatively, association of Rpd3 with Snt2 and Ecm5 may inactivate it. Consistent with the latter hypothesis, in Drosophila, association of Rpd3 with a complex containing the Ecm5 homolog Lid was reported to inhibit deacetylase activity (61).

We identified a second set of promoters where levels of Snt2 and Ecm5 increased after H$_2$O$_2$ stress (Fig. 4H and 5). While Rpd3 was enriched at these promoters by ChIP-qPCR, deletion of SNT2 or ECM5 did not affect Rpd3 association (Fig. 6D), suggesting that other proteins recruit Rpd3 to these targets. The Rpd3(L) complex regulates genes in response to diverse stresses (13–16), including oxidative stress (12), and may recruit Rpd3 to these regions. RNA-seq analysis identified 309 H$_2$O$_2$-enriched target genes that changed expression in wild-type cells in response to H$_2$O$_2$ and were either up- or downregulated when SNT2 was deleted (Fig. 7D), many of which function in oxidative stress response or oxidative metabolism (Fig. 7E).

While initially surprising that Snt2 might function as both an activator and a repressor, a number of other yeast transcriptional regulators, including Hap1 and Rap1, have been reported to have dual roles in transcription (58, 62, 63). The methods through which these proteins exert such opposing effects remain to be elucidated, but they may involve the cooperation of unique factors at different promoters. A subset of the H$_2$O$_2$-enriched promoters contained a motif similar to the known Pdr1/Pdr3 motif (Fig. 4I), and many of the Snt2/Ecm5 target genes regulated by Snt2 are also known to be targeted by Ste12 and Rap1 (Fig. 7H), suggesting that Pdr3, Pdr1, Ste12, and Rap1 may cooperate with Snt2 to regulate different gene sets.
support of this, the synthetic sickness of the ecm5Δ ask10Δ double mutant (17) suggests that Ecm5 function promotes oxidative stress tolerance, while Snt2 function ultimately decreases tolerance.

Because Snt2 and Ecm5 localize to genes that function in multiple pathways linked to the ESR, and not just to oxidative stress response genes, we hypothesized that rather than regulating only the oxidative stress pathway, Snt2 regulates stress pathways more broadly. In support of this, the snt2Δ strain was resistant to rapamycin (Fig. 8A), and rapamycin treatment promoted Snt2 and Ecm5 association with a subset of the H$_2$O$_2$, target promoters (Fig. 8C and D). A recent study reported that an snt2Δ strain was sensitive to histone overexpression (64). While the authors of that study propose that Snt2 functions in histone degradation, their results could also be explained if altered stress regulation in snt2Δ cells renders the cells more sensitive to the stress of histone overexpression.

The mechanism through which Snt2 and Ecm5 localization is regulated remains an exciting question and could involve stress-specific protein associations, posttranslational modifications, or regulation of subcellular distribution. Along these lines, our LC-MS analysis identified a phosphorylation site on serine 641 of Snt2 (Table 2). However, mutation of this serine to alanine or glutamate did not prevent ectopic Snt2 from rescuing the snt2Δ phenotype (data not shown), suggesting that this residue is not required for stress-mediated Snt2 regulation. It is noteworthy that a recent protein localization study reported that Ecm5 localized to the nucleus, while Snt2 localized to both the cytoplasm and nucleus and accumulated in the nucleus after nitrogen starvation (65). While this study did not find a change in Snt2 localization after H$_2$O$_2$ stress, we note that a higher concentration of H$_2$O$_2$ was used than in our study, possibly activating different stress pathways. Future studies aimed at providing insights into the mechanisms of Snt2 and Ecm5 regulation will be of considerable interest.

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