Comprehensive Transcriptional Analysis of the Oxidative Response in Yeast*§¶

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The oxidative stress response in Saccharomyces cerevisiae has been analyzed by parallel determination of mRNA levels and transcription rates for the entire genome. A mathematical algorithm has been adapted for a dynamic situation such as the response to stress, to calculate theoretical mRNA decay rates from the experimental data. Yeast genes have been grouped into 25 clusters according to mRNA level and transcription rate kinetics, and average mRNA decay rates have been calculated for each cluster. In most of the genes, changes in one or both experimentally determined parameters occur during the stress response. 24% of the genes are transcriptionally induced without an increase in mRNA levels. The lack of parallelism between the evolution of the mRNA amount and transcription rate predicts changes in mRNA stability during stress. Genes for ribosomal proteins and rRNA processing enzymes are abundant among those whose mRNAs are predicted to destabilize. The number of genes whose mRNAs are predicted to stabilize is lower, although some protein folding or proteasomal genes are among the latter. We have confirmed the mathematical predictions for several genes pertaining to different clusters by experimentally determining mRNA decay rates using the regulatable tetO promoter in transcriptional expression conditions not affected by the oxidative stress. This study indicates that the oxidative stress response in yeast cells is not only conditioned by gene transcription but also by the mRNA decay dynamics and that this complex response may be particularly relevant to explain the temporary down-regulation of protein synthesis occurring during stress.

Cells react against environmental stresses through multiple responses that occur at transcriptional and post-transcriptional levels to adapt themselves to the new conditions and counteract the possible macromolecular damage caused by the stress situation. Most systematic studies on such responses focus on changes in mRNA amounts (mRNA amount or concentration i.e. amount per cell), indicated as RA) caused by the environmental stress, using the DNA array technology. In the case of Saccharomyces cerevisiae, transcriptome analyses have been reported for a number of stresses including oxidative, osmotic, and nutritional ones in addition to heat shock (1, 2). Levels of a particular mRNA at a given time are the result of a balance between transcription rate (TR) and decay rate (commonly expressed as a half-life, or as a first-order kinetic constant of degradation, k0) (3). It is usually assumed that TR driven by specific transcriptional regulators plays the major role in the stress response. However, decay rate may change after the onset of an environmental stress and, in that instance, profiles of individual mRNAs may not directly reflect the corresponding TR profiles. We have developed a genomic run-on (GRO) methodology that allows quantifying TR and RA for each individual gene at a genomic scale (4). Moreover, mRNA half-lives can be obtained from TR and RA data under steady-state conditions. When applied to a nutritional shift from glucose to galactose, the GRO methodology showed that TR was the main determinant of RA, although some groups of genes were modulated at the mRNA decay level (4). More recently, we have developed a mathematical algorithm to determine mRNA half-life values from pointwise measurements of TR and RA in dynamic situations after the onset of an environmental stress when steady-state conditions cannot be assumed (3). Other groups have applied nuclear run-on approaches to culture cells. Fan et al. (5) determined RA and TR levels in human H1299 cells for about 1000 genes under non-stress and stress conditions, from which they inferred information on mRNA decay as well. Tennebaum et al. (6) developed the protocol “en masse run-on assay” in which the run-on profiling is combined with ribonomic profiling. Ribonomics is a term that defines the use of immunoprecipitated mRNP complexes to analyze the representation of individual mRNA species associated with a particular RNA-binding protein. The analyses of both kinds of data have led to the proposal that “post-transcriptional operons” (7) or “decay

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§1 The on-line version of this article (available at http://www.jbc.org) contains seven supplemental tables.

¶ The abbreviations used are: TR, transcription rate; pol I, RNA polymerase I; pol II, RNA polymerase II; GRO, genomic run-on; UTR, untranslated region; t-BOOH, tert-butyl hydroperoxide; RP, ribosomal proteins; GO, Gene Ontology.

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Decay of eukaryotic mRNA molecules may occur through different pathways (9–14). Two general mechanisms operate in *S. cerevisiae*: deadenylation of the 3′-poly(A) tail followed by 3′-5′ degradation of the mRNA by the exonuclease activity of the exosome or initial removal of the poly(A) tail followed by mRNA hydrolysis by the 5′-3′-exonuclease Xrn1.

*S. cerevisiae* is the first eukaryotic organism in which whole-genome studies have been done on mRNA stability in steady-state exponential growth conditions during glucose fermentation. In a first study (8), a temperature-sensitive RNA polymerase II (pol II) mutant was employed to switch off transcription and follow up mRNA levels at successive times for each transcript. In a second study (15), several chemical inhibitors of transcription were employed for the same purpose. Both studies showed that mRNA half-lives varied between a few minutes and more than 1 h and that mRNAs for proteins involved in ribosome biogenesis and rRNA processing were relatively unstable. A general correlation was also found between mRNA decay rates and the physiological function of the gene products (8). However, the experimental conditions employed for switching off transcription in the above studies caused a stress situation per se in yeast cells, which discards using a similar approach for analyzing mRNA kinetics after an experimentally induced stress (16).

Oxidative stress by external agents causes a transcriptional response in *S. cerevisiae* that involves a large number of genes (1), with the participation of the general stress response factor Msn2/4p and the oxidative stress-specific transcription factors Yap1 and Skn7 (17). Exposure of yeast cells to hydrogen peroxide results in increased levels of antioxidant proteins, heat shock proteins, components of the protein degradation machinery, and enzymes of the pentose phosphate pathway, which provides NADPH for some of the antioxidant systems (18, 19). In parallel, there is a down-regulation of the protein translation apparatus. This proteome pattern, although limited to about 20% of all expressed proteins in yeast cells, approximately parallels the transcriptome pattern (1), which indicates that the oxidative stress response mainly occurs through regulation of mRNA level. However, recent studies demonstrate that inhibition of protein synthesis occurring after an oxidative stress is not only caused by transcriptional down-regulation of the translation machinery but also by inhibition of translation initiation due to dissociation of ribosomes from mRNA and a slower rate of ribosomal runoff along mRNA molecules (20, 21). This illustrates the importance of the post-transcriptional level of regulation (22), which may also involve the regulation of mRNA translation efficiency by 3′-AU-rich elements, as is the case of the *MFA2* mRNA in *S. cerevisiae* (23). The oxidative stress response in the fission yeast *Schizosaccharomyces pombe* is in part regulated by the stabilization of atf1+ mRNA mediated by the RNA-binding protein Csx1 and Upf1 (24, 25). The latter is a component of the nonsense codon-mediated mRNA decay system. Atf1 is a transcription factor that coordinates the expression of many stress response genes and itself is under regulation of the Sty1 mitogen-activated protein kinase pathway (26).

The above studies in yeast cells leave unanswered the specific role of TR and mRNA decay in establishing new mRNA levels as a response to an oxidative stress. Here, we employ the GRO methodology to measure changes in TR and RA at a whole-genome level at different times after imposing an oxidative stress on yeast cells, and then we infer the evolution of mRNA half-lives after such a stress. Messenger RNA decay rates are validated for some genes whose expression becomes driven by the doxycycline-regulated *tet* promoter (27, 28), in conditions that do not cause an additional stress to yeast cells. Results indicate that for some functional groups of genes, changes in mRNA decay rates play an important role in the adaptation to oxidative stress.

**Experimental Procedures**

**Strains and Growth Conditions**—Wild type *S. cerevisiae* W303-1A (MATa ura3-1 ade2-1 leu2-3,112 trp-11 his3-11,15) was employed in the GRO experiments. MML830 is a derivative of the above strain by integration of EcoRV-linearized pCM244 (27). This plasmid codes for the tetR+ Ssn6 protein, which acts as repressor on tetO promoters after activation by doxycycline. The promoter-substitution cassette from plasmid pCM224 was employed for replacing the endogenous promoters of several genes in MML830 by the tetO promoter, as described in Ref. 28. The resulting strains are: MML863 (tetO2-HSP104), MML957 (tetO2-RP40), MML980 (tetO2-HSP42), and MML990 (tetO2-FIT3). In these strains, the gene expression driven by tetO2 is up-regulated in the absence of doxycycline, whereas the addition of the antibiotic represses it (27).

Cells were grown in YPD medium (1% yeast extract, 2% peptone, 2% glucose) at 28 °C. Doxycycline at 5 μg/ml was added for repressing genes under the tetO2 promoter. Oxidative stress conditions were created by the addition of t-butyl hydroperoxide (t-BOOH) at 0.1 mM. Experiments were initiated on exponential cultures that had been grown in such conditions for at least 10 generations, at concentrations of 1.5–2 × 107 cells/ml.

**Genomic Run-on**—Exponential cultures in 800 ml of YPD medium in 2-liter flasks shaken at 120 rpm were employed. Cell samples were taken at different times: t0 (exponential growth in YPD medium, just before t-BOOH addition), t1, t2, t3, t4, and t5, corresponding to 7, 16, 26, 41, and 71 min, respectively, after the application of the oxidative stress. At every sampling time, two different aliquots were taken. One of them was immediately processed to measure TR, according to the GRO protocol (see below). Cells from the second aliquot were recovered, washed with cold distilled water, frozen in liquid nitrogen, and stored at −80 °C until used for mRNA measurement.

The GRO protocol described previously (4) was used with the following modifications. Around 6 × 108 yeast cells were used to perform in vivo transcription. After spinning down cells, they were washed in cold water, and the cell pellet was resuspended in 900 μl of sterile cold water (final volume 950 μl). Then, the cell suspension was transferred to a fresh micro-centrifuge tube, 50 μl of 10% N-lauryl sarcosine sodium sulfate (sarkosyl) were added, and cells were incubated for 20 min on ice. After the permeabilization step, cells were recovered by low speed centrifugation, and the supernatant was removed. In vivo transcription was performed by resuspending cells in 120 μl of 2.5× transcription buffer (50 mM Tris-HCl, pH 7.7, 500 mM KCl, 80 mM MgCl2), 16 μl of AGC mix (10 mM each of CTP,
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ATP, and GTP), 6 µl of dithiothreitol (0.1 m), and 13 µl of 
[α-33P]UTP (3000 Ci/mmol, 10 µCi/µl). Cells were maintained on ice at all times. The final volume was adjusted to 300 µl with distilled water, and the mix was incubated for 5 min at 30 °C to allow transcription elongation. The reaction was stopped by adding 1 ml of cold distilled water to the mix. Cells were recovered by centrifugation to remove the non-incorporated radioactive nucleotide.

Total RNA was isolated using the Fast-Prep device (Bio101 Inc.) as described (4) except that acid phenol-chloroform (5:1) was used during cell breakage. Total extracted RNA was spectrophotometrically quantified. An aliquot was used for specific radioactivity determination in a scintillation counter. All the values for every gene (only one of the three replicates is allowed to be a removable outlier). Average cDNA values for each gene were finally corrected by the percentage of guanine residues in each probe-coding strand. Normalization between sampling points was made using the amount of mRNA/cell to give values of mRNA copies/cell for each gene in every time point. These values were used for cluster analysis and comparisons.

For normalizing the GRO hybridizations a different strategy was followed. First, we measured the total transcription per cell by using total dpm measured after extraction of the in vivo radioactively labeled RNA. The quantification of the extracted RNA allowed us to estimate the total transcription per cell. For each time point, we estimated TR (TR of pol I) by summing up the 16 signals from the specific probes (eight for 18 S and eight for 25 S regions) and TRII (TR of pol II) by summing up the whole set of signals (5950 pol II probes) in every hybridization. Since it is known that at t0 (exponential growth in YPD), the ratio between TR and TRII is about 2 (30) and that the estimated total transcription is, roughly, the sum of the contribution of these two polymerases, it is possible to determine the correction factor that fulfills these conditions. This factor was applied to the polymerase raw ratios for every sampling point. This allowed us to normalize the raw hybridization signals obtained for the RNA pol II probes. Statistical validation of replicates was performed as in cDNA values. After that, average TR values for each gene were finally corrected by the percentage of uracil residues present in each probe-coding strand. Again, the corrected average values were used for gene cluster analysis and other calculations.

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Nylon filters made using PCR-amplified whole open reading frame sequences as probes (29) were used as described (4) except that hybridizations were conducted during 40–48 h. Filters were exposed for 5–7 days to an imaging plate (BAS-MP, FujiFilm), which was read in a phosphorimaging scanner (FLA-3000, FujiFilm).

Measurement of RNA Levels—As mentioned above, a second cell aliquot was taken at each sampling time and immediately frozen with liquid nitrogen. After thawing the samples on ice, total RNA was isolated following the same procedure described for GRO samples except that RNA was precipitated once with 1 volume of 5 M LiCl, washed with 70% ethanol, resuspended in distilled water, and reprecipitated with 0.1 volume of 3 M NaAcO and 2 volumes of cold 96% ethanol. Again, RNA yield was measured spectrophotometrically. About 30–40 µg of DNA-digested total RNA were reverse-transcribed into cDNA as described (4) but using Invitrogen random hexamers for random priming. Hybridization was done in the same conditions as described for the GRO experiment except that labeled cDNA was at 5 × 106 dpm/ml and that filters were exposed for 1–2 days to an imaging plate.

Estimation of Total RNA and mRNA—To facilitate further normalizations, we estimated the RNA amount obtained from a fixed amount of cells along the experiment. Thus, five different cell aliquots were taken at each of the six sampling times (t0–t5) from a mock experiment. Total RNA was extracted, using the same protocol described previously, and quantified. Poly(A) mRNA was estimated using a dot-blot procedure as described (4). Using these data, we calculated the proportion of poly(A) mRNA per µg of total RNA and, thus, per cell at each of the time points (t0–t5, see Fig. 1).

Quantification of Hybridization Signals and Normalization Procedures—A total of six different nylon filters (one for each sampling time) were used. Filter combinations for the different hybridizations and replicates and image quantification were performed as described in Ref. 4. cDNA hybridizations were normalized within each experiment replicate by the global mean procedure. Correction factor to normalize between experiments was calculated from global mean values for the t0 sampling times. Reproducibility of the replicates was tested by the ArrayStat software (Imaging Research, Inc.), considering the data as independent and allowing the program to take a minimum number of valid replicates of 2 to calculate the mean values for every gene (only one of the three replicates is allowed to be a removable outlier). Average cDNA values for each gene were finally corrected by the percentage of guanine residues in each probe-coding strand. Normalization between sampling points was made using the amount of mRNA/cell to give values of mRNA copies/cell for each gene in every time point. These values were used for cluster analysis and comparisons.
every gene from its initial steady-state value ($t_0$) were summed up for all time points, and the sums for 4961 genes (those with values for six time points) were ranked.

**Clustering Procedures**—Changes in TR and RA, as well as mRNA stability estimates for all yeast genes, were evaluated by cluster analysis of normalized averaged values. For cluster analysis of the results, we used the Gene Expression Pattern Analysis Suite v 3.1 (GEPAS) included in the web server of CIPF Bioinformatic Unit, as described (4).

To test the potential enrichment in Gene Ontology (GO) categories in the different groupings obtained in this study, we used the FuncAssociate server, which uses a Monte Carlo simulation approach and accepts only significant GO categories according to their adjusted $p$ value (computed from the fraction of 1000 simulations under the null-hypothesis with the same or smaller $p$ value and after correction for multiple simultaneous tests). Only GO categories with an adjusted $p$ value below 0.05 were considered to be significant.

**Accession Numbers**—Gene Expression Omnibus (GEO) accession numbers for the whole experiment are GSE9645 (RA data) and GSE9663 (TR data).

**RESULTS**

**General Cell Responses to Oxidative Stress**—At certain non-lethal concentrations, hydroperoxides and other oxidants cause temporary growth arrest at the G1 stage of the *S. cerevisiae* cell cycle (31, 32). To avoid growth arrest, which could mask the direct effects of oxidative stress on general transcription and mRNA stability, we first tested a range of $t$-BOOH concentrations. We looked for conditions that did not affect exponential growth (when compared with control untreated cultures) but that still induced expression of three reporter genes (TRR1, TRX2, and HSP12), which have been shown to be induced by oxidative stress in previous studies (1, 33). We observed that 0.1 mM $t$-BOOH was the highest concentration of this oxidant that fulfilled such requirements. Therefore, all subsequent experiments were carried out with $t$-BOOH at 0.1 mM.

We then studied the general transcriptional responses of the cell after an oxidative stress. The total amount of poly(A) mRNA per cell decreased smoothly in the course of the experiment, whereas, as expected, cells continued proliferating (Fig. 1). Whole pol II transcription initially increased relative to time 0, an effect also observed after carbon source shift (4), to decrease at later times to levels similar to time 0. Relative TR levels did not decrease as much as relative RA levels, a fact that suggests a general mRNA destabilization following a moderate oxidative stress.

**Effect of Oxidative Stress on mRNA Stability**—The first-order degradation constants ($k_{dp}$, a measure of mRNA instability) can be estimated from the absolute values of TR and RA obtained by normalization of the GRO data. In a different study, we...
assumed steady-state conditions for most mRNAs and determined $k_D$ by division of RA by TR (4). However, in the current experiment, steady-state conditions are not expected to hold, at least during the first minutes of stress. Therefore, we employ here a mathematical approach based on the integration of the kinetic equations between two consecutive time points assuming a linear behavior of TR during the interval (3). Because $k_D$ is computed from absolute values of TR and RA, which have to be calculated by comparison with external data sets, the associated error might be enlarged by the mathematical manipulation.

Therefore, we used average profiles of gene groups to describe with more confidence the kinetic behavior of mRNA stability. The average data for each one of the 25 clusters are shown in Fig. 3A. Most clusters show an initial alteration of their $k_D$, but they tend to return to original values. Clusters 1–12, despite their differences, show a final $k_D$ higher than the initial one. Clusters 13–25 return to almost identical value to their initial $k_D$ with a slight trend to destabilization. Some clusters with barely detectable differences in TR+RA profiles show clearly distinct $k_D$ profiles (e.g. 9 versus 10, 23 versus 24), and some clusters with quite different TR+RA profiles (e.g. 7 versus 8) show very similar $k_D$ profiles, indicating that changes in mRNA stability are not easily deduced from RA+TR profiles during a dynamic situation. Profiles for clusters 1–5 but also for 14–17, 19, 20, and 23–25 show an initial decrease in $k_D$ followed by a fast recovery. The particular $k_D$ profiles are, however, different. For all these genes, it seems that stress causes a sudden mRNA stabilization whether being transcriptionally activated or not. The consequence of that stabilization is that RA increases more in the cases of up-regulation or decreases less for genes that are down-regulated. Clusters 6–12, the $k_D$ profile shows an opposite behavior with a sudden (clusters 6–9, 11, 12) or delayed (cluster 10) mRNA destabilization. It seems, therefore, that a main contribution to RA decrease in genes related to protein biosynthesis (mostly contained in clusters 6–7) is due to mRNA destabilization. Strikingly, some genes that showed a TR increase (clusters 8–11), including the ones for amino acid biosynthesis and those related to mitochondrial function, compensate or even down-regulate their RA by means of a hyperdestabilization of their mRNAs. Finally, only a small part of yeast genes (211, with no significant enrichment in specific GO categories) have approximately flat $k_D$ profiles (clusters 12 and 22), indicating that regulation of the mRNA stability is a general feature (>95%) of the oxidative stress response. The significance of $k_D$ as a regulatory mechanism is better seen when yeast genes are classified according to the magnitude of $k_D$ deviation from the steady-state value (Fig. 3B and supplemental Table S3). Some GO categories appear significantly enriched within the most affected
FIGURE 3. Predicted stability of mRNAs corresponding to the different gene clusters, after an oxidative stress. A, $k_d$ values are represented in the $y$ axis as a function of time (min) in the $x$ axis (t-BOOH added at time 0). The graphics represent the mean $k_d$ value corresponding to all the genes in the corresponding cluster in relative units referred to the mean $k_d$ value at time 0. Bars represent the standard error for each time point. $k_d$ scale is the same for all clusters, except for cluster 9.

B, histogram ranking total deviations from the initial $k_d$ during stress. The sum of differences (in absolute values) between the calculated $k_d$ and the initial steady-state $k_d$ (i.e. the $k_d$ at time 0) for all time points were calculated for the whole set of individual genes. The sum values were distributed in ranges, and the number of genes in each range interval is represented. Several GO categories related to ribosome biogenesis appear as statistically significant ($p$ value is shown) when genes with $\Sigma > 0.3$ are considered. In addition, the GO category “Organic Acid Transport” appears as significant when $\Sigma > 0.48$ is considered. Individual data can be seen in supplemental Table S3.
by changes in stability. Genes belonging to ribosome biogenesis categories and organic acid transport are specially controlled by changes in their mRNA stability. This is the first time that such a detailed analysis of mRNA stability has been done for a dynamic situation in any organism.

Several clusters display statistically significant negative values for \( k_D \) at certain time points (particularly during the first stages of the experiment). Negative \( k_D \) values (which make obviously no biological sense) are indicative of a final excess of RA over what could be expected from a linear evolution of TR between the values experimentally determined at the beginning and the end of the time interval. We are not aware of any artifact (such as a methodological bias or a release of mRNAs from a previously undetected pool) that could cause an eventual increase of RA consistently affecting only certain clusters at definite time points. Therefore, we believe that a possible explanation of the negative \( k_D \) values is that TR did not evolve linearly during that time interval but followed instead a pronouncedly convex trajectory, peaking between the experimentally determined values. Indeed, it has been argued that a transient TR peak is a fit transcriptional strategy for a fast transition to a new mRNA level after an environmental shift (3). Consequently, the analysis of the \( k_D \) profiles suggests an (experimentally undetected) transient TR peak between 0 and 7 min or/and between 7 and 16 min for some clusters (Fig. 3A). This hypothesis will be investigated in the future.

**Experimental Determination of mRNA Decay Rate for Some Representative Genes**—We employed an experimental approach to confirm the kinetics of mRNA decay for some genes after application of an oxidative stress. For this purpose, promoters of the corresponding genes were substituted by the doxycycline-regulatable \( \text{tetO} \) promoter, and mRNA decay rates were determined in the resulting strains by measuring mRNA signal levels in Northern blots at different times after the addition of doxycycline (see “Experimental Procedures”). Previously, we had shown that down-regulation of \( \text{tet} \) promoters using the activator (tTA)-repressor (tetR-Ssn6) dual system occurs very shortly after the addition of doxycycline (27). For each experiment, mRNA decay rate was determined just before the addition of t-BOOH and at two different times after the addition of the oxidant, which were selected based on the kinetics of decay according the \( k_D \) values predicted by the mathematical algorithm for each particular gene (supplemental Table S1).

Heat shock genes \( \text{HSP42} \) and \( \text{HSP104} \) are in clusters 15 and 16 respectively (supplemental Table S2). In both cases, transcript level and TR, as determined in the GRO experiment, increase transitorily after the addition of the oxidant and decrease to near original values at later times (supplemental Table S1). Increase of TR precedes that of mRNA, and it is mathematically predicted for clusters 15 and 16 that mRNA molecules are transitorily stabilized (lower \( k_D \)) at initial times after the onset of stress (Fig. 3A). Differences between genes in clusters 15 and 16 basically rely upon the fact that TR is more intensely up-regulated in the case of cluster 16 (Fig. 2). We confirmed by Northern analyses that \( \text{HSP42} \) mRNA level peaked around min 20 after the addition of the oxidant (Fig. 4A) following kinetics similar to that in the GRO experiment. The

![FIGURE 4. Experimental determination of mRNA half-lives before and after the addition of t-BOOH.](image-url)

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Strains expressed \( \text{HSP42} \) (A, strain MML980), \( \text{HSP104} \) (B, MML863), \( \text{RRP40} \) (C, MML957), or \( \text{FIT3} \) (D, MML990) under the control of the \( \text{tetO} \) promoter. Doxycycline (5 \( \mu \)g/ml) was added at time 0 to exponentially growing cultures of the corresponding strains at 28 °C that had not been exposed to t-BOOH or 10, 20, or 60 min after the addition of t-BOOH. In each case, aliquots were taken at time 0 and at successive times after the addition of doxycycline, for total mRNA isolation and determination of levels of the corresponding mRNA by Northern analysis. Graphics represent the evolution of experimentally determined relative RA in log scale as a function of time for a representative experiment. Mean half-life values (\( t_{1/2} \) plus standard deviation for a total of three independent experiments) are also indicated. To determine \( t_{1/2} \) values, linear regression of experimental data (as represented in the figure) was calculated, exclusively considering only the initial points for which linearity was maintained. The upper panels show Northern analyses of RA expressed under the respective own promoters in wild type (W303-1A) cells growing exponentially after adding t-BOOH at time 0. U1 RNA is included as loading control.
half-life of the HSP42 mRNA was determined before the stress and at min 20 and 60 after it (Fig. 4A). As expected (34), initial mRNA decay rates followed first-order kinetics, and half-lives were calculated from the slopes of the log curves. From a half-life value of 14.3 min during exponential growth, this increased to 18.6 min after 20 min of applying the stress, to decrease to 7.6 min after 60 min in stress conditions (Fig. 4A). That is, HSP42 mRNA became initially stabilized, and it was destabilized at later times relative to unstressed cells. Both the initial stabilization and the final destabilization confirmed the mathematical predictions (Fig. 3A, cluster 16). A similar picture was experimentally determined for HSP104 mRNA. Thus, there was a temporary up-regulation of mRNA levels as determined by Northern analysis and an initial transitory increase of mRNA half-life (Fig. 4B). In this case, final decay rate 60 min after the stress was close to that of unstressed cells.

Members of clusters 19 and 20 experience a much higher and rapid increase in mRNA levels than in TR upon the oxidative stress (Fig. 2), which consequently predicts a temporary stabilization of mRNAs (Fig. 3A). We tested this prediction for FIT3, a member of cluster 20 that shows an intense transitory up-regulation of mRNA levels in response to peroxide (Fig. 4C). The FIT3 product is a cell wall mannoprotein that binds siderophore-iron chelates, therefore facilitating iron uptake (35). FIT3 mRNA half-life almost doubled 10 min after the stress relative to unstressed cells, whereas at 60 min, the mRNA half-life value had approximately returned to the situation in untreated cells (Fig. 4C). This confirmed that initial stabilization of FIT3 mRNA contributed to the transitory up-regulation of mRNA levels.

The mathematical model employed in this work predicts a temporary destabilization of mRNAs of genes in cluster 6 (Fig. 3A), which includes many genes for ribosomal proteins (RP) and for rRNA-processing proteins (supplemental Table S2). We tested this prediction for RRP40 mRNA, whose levels and TR decrease after the onset of stress as determined by GRO, in parallel to a predicted transitory destabilization of mRNA molecules (supplemental Table S1). Northern analyses confirmed the down-regulation of RRP40 mRNA levels upon oxidative stress (Fig. 4D). Decay kinetics showed that RRP40 mRNA dramatically destabilized after imposing the stress, from an initial half-life value of 8.7 min in unstressed cells to 4.7 and 3.4 min after 10 and 60 min of t-BOOH addition, respectively (Fig. 4D). Thus, the experimental work using the regulatable tet promoter system confirmed the prediction of a stress-induced mRNA destabilization.

Correlation between Transcription Parameters and Gene Function—mRNA level profiles in response to environmental stresses tend to correlate among functionally related genes (36, 37). Thus, levels of mRNA for RP and for rRNA-processing proteins decrease upon oxidative stress (1). We extended these previous analyses to the other two mRNA-related parameters (TR, decay rate) from the GRO kinetic data (supplemental Table S1) in the oxidative stress response. 122 RP genes (most of them at cluster 7 in Fig. 2) for which time course data existed from at least two independent experiments were selected. Fig. 5 shows the global results for the whole group of genes (mean values), and supplemental Table S4 lists the results for the individual genes. Levels of mRNAs decrease steadily during the first 40 min of stress, maintaining afterward a constant value at less than 30% relative to non-stressed conditions, whereas TR decreased transitorily between 20 and 30 min to then recover afterward to control levels (Fig. 5A). This correlates with the predicted sustained destabilization of mRNAs (Fig. 5B). Similar general profiles were obtained for 135 genes in the “rRNA Processing” GO category (mainly in cluster 6 in Fig. 2), listed in supplemental Table S5. However, TR for rRNA-processing genes did not recover along the experimental period (Fig. 5A). mRNAs for this GO category were predicted to destabilize upon the stress, although at sustained lower levels than for RP mRNAs (Fig. 5B). The predicted destabilization correlates with the above results for RRP40 mRNA (Fig. 4C). Interestingly, a modest initial up-regulation of TR was observed after oxidant addition for both RP and rRNA-processing genes (Fig. 5A).

The environmental stress response causes up-regulation of the mRNAs for proteasome subunits (1, 37). In the particular case of the oxidative stress response, it has also been shown that many proteasome subunit proteins are induced (18). We analyzed the profiles of 14 genes included in the GO category “Proteasome Core Complex” (Fig. 6 and supplemental Table S6). They are distributed among eight different clusters in Fig. 2, clusters 12 and 24 containing three genes each. In general, TR for proteasome core complex genes displayed a rapid and transitory induction to decrease below initial levels at later times (Fig. 6). This increase preceded that of mRNA levels. TR for most mRNAs coding for the proteasome core complex is coordinately up-regulated immediately after the onset of the oxidative stress, and this is followed by late repression. This by itself can explain the delayed parallel changes in mRNA levels. The coordinated behavior of TR for core proteasome genes can be related to the role of the Rpn4 transcription factor as regulator of expression of most proteasome yeast genes (38). However,
the RA profiles are much more variable, indicating that posttranscriptional mechanisms operate differentially on them. Analysis of genes in the GO category "Proteasome Regulatory Complex" resulted in similar patterns to those of the core complex (data not shown).

The GO category "Oxidoreductase Activity" is significantly overrepresented in cluster 8 (Fig. 2), and a number of additional members of the category are in clusters 9–12 (supplemental Table S7). As indicated above, clusters 8–12 show an increase in TR not accompanied by RA increase. To further analyze this category, we divided oxidoreductase activity genes into those in clusters 8–12 and those in the rest of clusters (supplemental Table S7). When mean values of TR and RA are represented separately for both subgroups (Fig. 7), two different behaviors can be observed: those genes that expectedly show a parallelism between TR and RA, with a modest increase in TR preceding that of RA (clusters 1–7 and 13–25), and those that show a higher increase in TR while keeping RA basically constant, which predicts a destabilization of mRNAs (clusters 8–12). Promoter and 3'-untranslated region sequence in silico analysis did not evidence statistically significant enrichment of specific sequences in any of both groups of genes. Nevertheless, it is remarkable that up to one-third of oxidoreductase genes in clusters 8–12 (which would have not been detected in genomic studies strictly based on RA analyses) contain Yap1-recognizable sequences in their promoters and about 40% contain STRE sequences recognized by the Msn2/Msn4 factors. The presence of such sequences explains the TR increase upon the stress application.

DISCUSSION
In this work, we have extended previous studies on the effect of oxidative stress on mRNA amounts at the whole transcriptome level (1, 2). Thus, we have carried TR analyses and consequently predicted effects of the stress on mRNA stability. Conditions were employed in which growth was not affected after the addition of the oxidant, thereby discarding transcriptome-level effects caused by growth rate changes. Globally, pol II-mediated transcription rate did not change significantly during the examined time interval, except for a brief up-regulation at the onset of stress. However, specific sets of genes displayed higher TR values, whereas others had their TR diminished in the oxidative conditions employed.
In most of the 4757 genes (about 80% of the yeast genome) for which complete TR and RA data were obtained in this study, changes in one or both parameters occurred during the oxidative stress response. It is interesting to note that our method detects changes in gene transcription due to oxidative stress that could not be detected by conventional analyses. Thus, 1147 genes belonging to clusters 8–12 do not show an increase in RA, the usual parameter evaluated by most DNA chip analyses, but they did show an increase in TR. This means that they suffer a general destabilization in their mRNA along the time course, as shown in Fig. 3A. Therefore, we have extended the group of transcriptionally induced genes by oxidative stress by more than 2-fold: 1968 genes showing TR increase versus 821 that only show increase in RA. Because changes in TR are the primary consequence of the action of transcription factors, we propose that the analyses of TR changes by the GRO methodology will improve the searches (e.g. Ref. 39) for genes belonging to the same regulon. On the other hand, 768 genes in cluster 7 (including most RP genes, Fig. 5A) that show a significant RA down-regulation over the time course only show a minute and fluctuating TR change. Both examples illustrate the influence of mRNA stability in gene regulation and the power of our approach to detect it. Certainly, for a significant number of genes, changes occurred at modest levels. This is the case for cluster 5 (439 genes), which is characterized by a modest down-regulation (less than 50% decrease when compared with time 0) of TR and RA and is enriched in genes involved in protein secretion. Clusters 11, 12, 22, 23, and 24 also display rather constant values of TR and RA, without being enriched in particular functional categories. In total, less than one-fourth of the yeast genome maintain rather stable values (less than 2-fold changes) of both TR and RA at the oxidative stress conditions applied in this study. Clusters 6 and 7 (1391 genes) exhibit a large decrease of RA accompanied by TR down-regulation, which is more dramatic in the genes of cluster 6. The two clusters include most of the genes implicated in ribosome structure and biogenesis (RP and rRNA-processing enzymes), and this reflects the inhibition of protein synthesis after application of oxidative stress (21). However, TR inhibition alone is not sufficient to explain the decrease in RA in clusters 6 and 7 genes. The mathematical model employed here predicts a significant destabilization of mRNA molecules upon stress, which would be co-responsible for such decrease. When genes for RP and for rRNA-processing enzymes are analyzed separately from other genes in the clusters to which they pertain, an even more dramatic increase in mRNA decay rate is predicted for both groups, and we have experimentally confirmed it for RRP40. In ribosome-related genes, mRNA destabilization is maintained for at least 60 min after the stress. We may conclude that the down-regulation in the expression of ribosome-related genes and the subsequent inhibition of protein synthesis after an oxidative stress result from additive contributions of inhibition of transcription and increased decay rate of the respective mRNAs.

Most of the clusters from 8 to 25 display an immediate induction of TR upon the oxidative stress. For some of them, RA increase is delayed a few minutes relative to TR, as would be expected whether changes in RA were a direct consequence of TR changes (3). However, there is the exception of clusters 8–12, already commented. Clusters 13, 15, and 16 are representative of those showing an approximate parallelism between TR and RA up-regulation. These three clusters are respectively enriched in amino acid metabolism, trehalose synthesis, and sulfur metabolism genes. Activation of trehalose synthesis as a protective mechanism in response to different environmental stresses, including oxidative stress, has already been described (1). Induction of a number of pathways of amino acid biosynthesis as a response to the moderate stress conditions employed here may be an adaptive strategy to prepare cells for protein synthesis recovery. Particularly relevant is the strong induction of the biosynthetic pathway for sulfur amino acids observed in this study. Such induction has not been reported in other studies on oxidative stress responses where a higher oxidant concentration was employed (1). However, induction of this pathway by cadmium (40) and arsenite (41) has been described using genomic and proteomic approaches. Up-regulation of the sulfur amino acid pathway would thus result in higher levels of glutathione (which requires cysteine for its synthesis) needed for the redox response against oxidant conditions. Glutathione is the substrate for glutaredoxins, a group of thiol oxidoreductases participating in the oxidative stress response (42, 43). These, together with other oxidoreductases and additional enzymes detoxifying reactive oxygen species, were induced in our study. They are distributed among different clusters, especially in clusters 8 and 9.

Even for the genes (or groups of genes) with the parallel kinetics of transitory up-regulation of TR and RA, changes in mRNA decay rate may influence the response pattern. The k_D values calculated with the new algorithm may be meaningful except for cases of very fast and transitory TR responses (usually restricted to the first minutes of stress) for which the assumed linear evolution of TR in between experimental points does not hold, producing artificial negative values of k_D. Nevertheless, negative k_D values may help to identify genes that respond to stress through short-lived abrupt TR peaks, which may be studied in the future by means of a more frequent sampling. In any case, the k_D profiles calculated by the new procedure clearly show that changes in stability are characteristic of most of the mRNAs after the oxidative stress and that the main part of their change is transitory, restricted to the first 20 min. We have studied in detail some interesting examples, such as core proteasome genes, whose mRNA levels decay at late times more slowly than TR, suggesting that mRNA stabilization is part of the stress recovery response. It is also the case for a number of genes in the "Protein Folding" GO category (data not shown in detail). We have confirmed this prediction for two heat shock proteins in this category, HSP42 and HSP104. In both cases, temporary stabilization may contribute to the fact that up-regulation of RA lasts longer than that of TR. We have tested the mathematical results for a total of four examples of genes corresponding to different clusters, including the mentioned HSP42 and HSP104 genes, and all of them qualitatively coincide with predictions (Fig. 4). In clusters 15 and 16, although the predicted k_D values at short times are not valid, the stabilization of the mRNA is corroborated. For some functional categories, such as ribosome biogenesis and organic acid transporter (Fig. 3B) and ribosomal proteins (Fig. 5), the influence of
mRNA stability is especially important as already suggested (8, 15). Our study shows, for the first time, the extension of such a mechanism for a dynamic response in a cell.

The behavior of oxidoreductase-coding genes in clusters 8–12 deserves special attention since temporary increase in TR is not accompanied by significant changes in RA, predicting temporary mRNA destabilization upon oxidative stress. A similar situation occurs with amino acid biosynthetic genes, which are also overrepresented in clusters 8–12. Using in silico tools, we have not been able to detect specific sequences in promoter or terminator regions of genes for oxidoreductases, which could explain such differences between TR and RA. This leaves open the question on the molecular determinants of the lack of parallelism between both parameters. Concerning the significance of the futile transcription of a subpopulation of mRNAs to be immediately degraded, we can hypothesize that such genes may also respond to different stresses during which they would require high transcript levels. Therefore, the promoter elements of each particular gene would be responsible for a common TR response upon the diverse stresses, but later modulation of RA would accommodate the response to each specific stress. Testing this hypothesis will require to extend this type of studies to other stresses.

In summary, analysis of TR upon a stress gives significantly more information than simply measuring changes in RA. Importantly, determining both parameters allows making inferences on how mRNA stability influences the oxidative stress response. In addition, by using a genetic system for ectopic regulation of expression of particular genes under stress in conditions not additionally influencing the cell physiology, we have been able to confirm that changes in mRNA decay rates are indeed part of the oxidative stress response for certain groups of genes. This raises the interest in searching for a mechanistic connection between oxidative stress and decay of mRNA molecules in S. cerevisiae, as occurs in fission yeast through the Csx1 and Upf1 proteins and the Atf1-mediated stress response (24, 25).

REFERENCES
1. Gasch, A. P., Spellman, P. T., Kao, C. M., Carmel-Harel, O., Eisen, M. B., Storz, G., Botstein, D., and Brown, P. O. (2000) Mol. Cell. Biol. 11, 4241–4257
2. Causton, H. C., Ren, B., Koh, S. S., Harbison, C. T., Kanin, E., Jennings, E. G., Lee, T. I., True, H. L., Lander, E. S., and Young, R. A. (2001) Mol. Biol. Cell 12, 323–337
3. Pérez-Ortín, J. E., Alepuz, P. M., and Moreno, J. (2007) Trends Genet. 23, 260–267
4. García-Martínez, J., Aranda, A., and Pérez-Ortín, J. E. (2004) Mol. Cell 15, 303–313
5. Fan, J., Yang, X., Wang, W., Wood, W. H., III, Becker, K. G., and Gorospe, M. (2002) Proc. Natl. Acad. Sci. U. S. A. 99, 10611–10616
6. Tenenbaum, S. A., Carson, C. C., Atasoy, U., and Keene, J. D. (2003) Gene (Amst.) 317, 79–87
7. Keene, J. D., and Tenenbaum, S. A. (2002) Mol. Cell 9, 1161–1167
8. Wang, Y., Liu, C. L., Storey, J. D., Tishbarami, R. I., Herschlag, D., and Brown, P. O. (2002) Proc. Natl. Acad. Sci. U. S. A. 99, 5860–5865
9. Wilusz, C. J., Wormington, M., and Peltz, S. W. (2001) Nat. Rev. Mol. Cell Biol. 2, 237–246
10. Chen, C. Y., Gherzi, R., Ong, S. E., Chan, E. L., Rajmakers, R., Pruijn, G. J., Stoecklin, G., Moroni, C., Mann, M., and Karin, M. (2001) Cell 107, 451–464
11. Parker, R., and Song, H. (2004) Nat. Struct. Mol. Biol. 11, 121–127
12. Maquat, L. E. (2004) Nat. Rev. Mol. Cell. Biol. 5, 89–99
13. Newbury, S. F. (2006) Biochem. Soc. Trans. 34, 30–34
14. Simon, E., Camier, S., and Séraphin, B. (2006) Trends Biochem. Sci. 31, 241–243
15. Grigull, J., Minaimneh, S., Pootoolal, J., Robinson, M. D., and Hughes, T. R. (2004) Mol. Cell Biol. 24, 5534–5547
16. Pérez-Ortín, J. E. (2008) Brief. Funct. Genomics Proteomics 6, 282–291
17. Toledano, M. B., Denuay, A., Bittau, B., Spector, D., and Azevedo, D. (2003) in Topics in Current Genetics (Hohmann, S., and Mager, P. W. H., eds) Vol. 1, pp. 241–303, Springer-Verlag, Berlin
18. Godon, C., Lagniel, G., Lee, J., Buhrer, J.-M., Kieffer, S., Perrot, M., Boucherie, H., Toledano, M. B., and Labarre, J. (1998) J. Biol. Chem. 273, 22480–22489
19. Lee, J., Godon, C., Lagniel, G., Spector, D., Garin, J., Labarre, J., and Toledano, M. B. (1999) J. Biol. Chem. 274, 16040–16046
20. Swaminathan, S., Masek, T., Molin, C., Pospisik, M., and Sunnerhagen, P. (2006) Mol. Cell Biol. 17, 1472–1482
21. Shenton, D., Smirnova, J. B., Selley, J. N., Carroll, K., Hubbard, S. J., Pavitt, G. D., Ashe, M. P., and Grant, C. M. (2006) J. Biol. Chem. 281, 29011–29021
22. Sunnerhagen, P. (2007) Mol. Genet. Genomics 277, 341–355
23. Vassudev, S., Garneau, N., Khounh, D. T., and Reltz, S. W. (2005) Mol. Cell Biol. 25, 9753–9763
24. Rodríguez-Gabriel, M. A., Burns, G., McDonald, W. H., Martin, V., Yates, J. R., III, Bühler, I., and Russell, P. (2003) EMBO J. 22, 6526–6526
25. Rodríguez-Gabriel, M. A., Watt, S., Bühler, J., and Russell, P. (2006) Mol. Cell Biol. 26, 6347–6356
26. Shiozaki, K., and Russell, P. (1996) Genes Dev. 10, 2276–2288
27. Belli, G., Gari, E., Piedrafita, L., Aldea, M., and Herrero, E. (1998) Nucleic Acids Res. 26, 942–947
28. Belli, G., Gari, E., Aldea, M., and Herrero, E. (1998) Yeast 14, 1127–1138
29. Alberola, T. M., García-Martínez, J., Antúnez, O., Viladevil, L., Barceló, A., Ariño, J., and Pérez-Ortín, J. E. (2004) Int. Microbiol. 7, 199–206
30. Warner, J. R. (1999) Trends Biochem. Sci. 24, 437–440
31. Lee, J., Romeo, A., and Kosman, D. J. (1996) J. Biol. Chem. 271, 24885–24893
32. Wanke, V., Accorsi, K., Porro, D., Espósito, F., Russo, T., and Vanoni, M. (1999) Mol. Microbiol. 32, 753–764
33. Ocon-Garrido, E., and Grant, C. M. (2002) Mol. Microbiol. 43, 993–1003
34. Hargrove, J. L., Hulse, M. G., and Beale, E. G. (1991) BioEssays 13, 667–674
35. Protchenko, O., Ferea, T., Rashford, J., Tiedeman, J., Brown, P. O., Botstein, D., and Philpott, C. C. (2001) J. Biol. Chem. 276, 49242–49250
36. Wu, L. F., Hughes, T. R., Davierwala, A. P., Robinson, M. D., Stoughton, R., and Altschuler, S. J. (2002) Nat. Genet. 31, 255–265
37. Gasch, A. P. (2003) in Topics in Current Genetics (Hohmann, S., and Mager, P. W. H., eds) Vol. 1, pp. 11–70, Springer-Verlag, Berlin
38. Mannhaupt, G., Schnall, R., Karpov, V., Vetter, I., and Feldmann, H. (1999) FEBS Lett. 450, 27–34
39. Buissermaker, H. J., Hao, L., and Siggia, E. D. (2001) Nat. Genet. 27, 167–171
40. Thorsen, M., Lagniel, G., Kristiansson, E., Junot, C., Nerman, O., Labarre, J., and Tamás, M. J. (2007) Physiol. Genomics 30, 35–43
41. Toledano, M. B., Kumar, C., Le Moan, N., Spector, D., and Tacket, C. (2007) FEBS Lett. 581, 3598–3607
42. Herrero, E., Ros, J., Belli, G., and Cabaiscol, E. (2008) Biochim. Biophys. Acta 10.1016/j.bbaben.2007.12.004