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Genome-wide single-cell-level screen for protein abundance and localization changes in response to DNA damage in S. cerevisiae

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ABSTRACT

An effective response to DNA damaging agents involves modulating numerous facets of cellular homeostasis in addition to DNA repair and cell-cycle checkpoint pathways. Fluorescence microscopy-based imaging offers the opportunity to simultaneously interrogate changes in both protein level and subcellular localization in response to DNA damaging agents at the single-cell level. We report here results from screening the yeast Green Fluorescent Protein (GFP)-fusion library to investigate global cellular protein reorganization on exposure to the alkylating agent methyl methanesulfonate (MMS). Broad groups of induced, repressed, nucleus- and cytoplasm-enriched proteins were identified. Gene Ontology and interactome analyses revealed the underlying cellular processes. Transcription factor (TF) analysis identified principal regulators of the response, and targets of all major stress-responsive TFS were enriched amongst the induced proteins. An unexpected partitioning of biological function according to the number of TFs targeting individual genes was revealed. Finally, differential modulation of ribosomal proteins depending on methyl methanesulfonate dose was shown to correlate with cell growth and with the translocation of the Sfp1 TF. We conclude that cellular responses can navigate different routes according to the extent of damage, relying on both expression and localization changes of specific proteins.

INTRODUCTION

Several DNA repair and cell-cycle checkpoint pathways have evolved to cope with damage to the genome that can arise from endogenous and exogenous sources (1,2). It is well established that effective cellular responses to DNA damaging agents involve not only modulation of canonical DNA repair and cell-cycle regulation proteins but also modulation of a large number of seemingly unrelated cellular processes (3). Previous studies identified a number of these pathways using ‘transcriptional profiling’ and ‘genomic phenotyping’ in Saccharomyces cerevisiae (4–9). Transcriptional profiling quantified global changes in mRNA levels in response to genotoxic stress using microarrays, whereas genomic phenotyping investigated the sensitivity to DNA damaging agents for almost 6000 S. cerevisiae strains, each deficient in a single gene product. A consensus emerged from these and other studies (9,10) that a general shutdown of protein synthesis occurs under conditions of DNA damage because the transcription of ribosomal protein (RP) genes is repressed under conditions of genotoxic stress. Later studies demonstrated a concomitant preferential translation of specific damage-responsive proteins under conditions of genotoxic stress (11–13), arguing for translational regulation playing a role in the cellular response to DNA damaging agents. Finally, a number of post-transcriptional protein modifications are known to orchestrate the DNA damage response, including phosphorylation, ubiquitylation and sumoylation (1,14).

To probe response pathways at the single-cell level, we developed a quantitative high-throughput fluorescence imaging approach to assess not only changes in protein levels but also changes in nuclear versus cytoplasmic localization in response to the DNA damaging agent methyl methanesulfonate (MMS). This assay was performed for >4000 S. cerevisiae strains expressing individual GFP-tagged fusion-proteins, representing nearly 70% of the yeast proteome (15). Importantly, fusion proteins in this library are expressed from their native promoters, thereby closely reflecting the response of their corresponding genes on MMS exposure.
Previously, this library was used to study genome-wide protein localization using fluorescence microscopy (15) and overall expression levels using flow cytometry (16). Here, we use the library to identify proteins whose expression level is induced or repressed in response to MMS, as well as proteins that concentrate in either the nuclear or cytoplasmic compartment in response to MMS. Analysis of Gene Ontology (GO) and protein–protein interaction networks reveal damage-induced changes in levels and/or localizations for proteins and complexes involved in chromatin remodeling, mRNA processing, RNA polymerase II transcription, proteolysis, ribosome biogenesis, metabolism, lipid synthesis, plus a number of other pathways, in addition to canonical DNA repair and cell-cycle regulation. Further, we characterize the transcription factor (TF) networks linked to changes in protein abundance, revealing a differential regulation of metabolic versus DNA-related processes. Finally, we further investigate the unexpected induction response of RPs, finding a differential response depending on the extent of damage, cellular growth rate and nuclear-to-cytoplasmic translocation of the TF Sfp1 that targets the RP genes (17).

MATERIALS AND METHODS

Cell growth and culture conditions

We used the budding yeast GFP fusion library developed by Huh and colleagues (15). The haploid parent yeast strain was ATCC 201388: MAT\textit{his3A1leu2A0met15D0ura3D0}. This strain is called wild-type (WT) throughout the manuscript. Cells were cultured in minimal SD medium (MP Biomedicals) supplemented with amino acids His, Leu, Ura and Met. Cells were grown at 30°C. Cells were cultured to stationary phase for 3 days and then diluted in fresh medium and allowed to grow overnight in triplicate cultures. Log-phase cultures were incubated in growth medium with or without 0.02% MMS for 3 h. Details of plate preparation for High Content Imaging are presented in the Supplementary Information. For the initial purposes of identifying a better fixation method, either one of two protocols adapted from previous studies was followed (18,19) (Supplementary Information, Supplementary Figures S1–S5). For all subsequent experiments, the second fixation method (called Fix 2) was followed, as it caused lesser loss of GFP fluorescence and also did not introduce any additional autofluorescence (Supplementary Figures S3–S5). Plates for the screen were prepared in triplicate on a Tecan liquid handling robot (Mannedorf, Switzerland) running on EVOWare software.

Cell processing for flow cytometry and fluorescence microscopy

Cells were treated with 1 μg/ml 4’,6-diamidino-2-phenylindole (DAPI) and 2.5 μg/ml Concanavalin A-Alexa647 (both from Invitrogen Life Technologies) for 30 min to stain DNA and the cell-wall, respectively. Cells used for flow cytometry were resuspended in phosphate buffered saline, whereas cells used for imaging were mounted on concanavalin A-coated 96-well plates to allow adhesion of the cells onto the bottom of the well. Excess cells were washed off and the remaining cells were mounted in 30% glycerol in phosphate buffered saline. Samples were prepared in triplicate for both control untreated and MMS-treated samples.

Flow cytometry and fluorescence microscopy

Flow cytometry was performed on an Accuri C6 Flow Cytometer (Accuri Cytometers Inc., Ann Arbor, MI), unless otherwise mentioned. Imaging was performed on a Cellomics Arrayscan VTi (Thermo Fisher Scientific, Pittsburgh, PA), using the XF93 filter to image GFP, DAPI and Alexa 647 fluorescence. A 40× 0.75 Numerical aperture (N.A.) air objective was used for imaging. High-resolution imaging was performed on an Observer Z1 microscope (Carl Zeiss, Jena, Germany) with a 100× 1.4 N.A. oil immersion objective.

Image analysis

Image analysis was performed using MATLAB (MathWorks Inc., Natick, MA) using custom-written routines for the detection of cellular boundaries from images of the cell-wall stained with Alexa 647 conjugated Concanavalin A. Clusters of cells were eliminated from calculations because nuclear versus cytoplasmic localizations could not be correctly computed for these. DAPI-stained images of DNA were used for computing nuclear positions by thresholding out the nuclei after elimination of cells, which showed uniform DAPI staining by setting cutoff conditions of intensity and area. Masks of nuclei thus obtained were used as a mask on the GFP image to compute GFP fluorescence levels in the nucleus. The rest of the cell was treated as cytoplasm, enabling the computation of nuclear to cytoplasmic ratios. In our approach, the mean GFP intensities are evaluated in the whole cell, nuclear and cytoplasmic masks. This is distinct from a previous work that used an expanded nuclear mask to evaluate cytoplasmic fluorescence (20). All raw images, data analysis programs and single cell level files associated with this study can be accessed at: http://yeastgfpscreen.mit.edu/.

Data analysis

Experiments were performed in triplicate and the mean of each replicate was calculated. The WT control was used to estimate the autofluorescence level, which was subtracted from all the measured strains in that experiment. WT controls were present in every plate. We found that for many low-expressing strains when the measured intensity is close to autofluorescence, there can be spuriously high estimates of fold-changes. Such strains with fluorescence levels close to autofluorescence in the control samples were retained in the analysis because the expression of a protein may be turned on by MMS treatment. To avoid spuriously high fold-changes, we added a constant value, c (equal to the width of the autofluorescence histogram), to every measured value. Although this leads us to underestimate the fold-changes, this ensures that only substantial changes are scored as true responders. Thus, fold-change for an experimental strain is calculated as, $f = \frac{[I_{\text{MMS}} - (I_{\text{WT}} - c)]}{[I_{\text{Control}} - (I_{\text{WT}} - c)]}$, where $I$ denotes mean
intensity from triplicate samples, and the subscripts denote the conditions and strains. The autofluorescence estimated from the WT strain [as also done in a previous study (21)] is only a ballpark value, and there can be a strain-to-strain variability of true autofluorescence (16). This approach ensures that changes in expression due solely to differences in autofluorescence are not scored as responders. However, because experiments are performed in triplicate, statistical confidence can be attributed even to relatively small changes when they pass the thresholds used. The individual means, obtained from the three experiments, were compared by Student’s t-tests at $P \leq 0.05$ and $f \geq 1.5$ (for induced proteins) or $f \leq 0.75$ (for repressed proteins). For nuclear-to-cytoplasmic ratio (NCR) changes by t-tests, $P \leq 0.05$ and $|\Delta \text{NCR}| \geq 0.1$ were used. $\Delta \text{NCR} = \text{NCR}_{\text{Control}} - \text{NCR}_{\text{MMS}}$, and a negative sign indicates nuclear enrichment, whereas a positive sign indicates cytoplasmic enrichment. Typically hundreds to thousands of cells were measured for each strain. If for any particular well the cell count was $\leq 50$, that well was eliminated from all calculations.

Protein intensity distributions over cells are often non-Gaussian. However, the means arising from repeated measurements of such distributions are normally distributed, and thus Student’s t-test can be used for comparing the means obtained from such measurements. This assumption of normality does not hold true when comparing the underlying distributions of protein intensities over cells, in which case a non-parametric test is preferable. We used the Kolmogorov–Smirnov (KS) test for determining responders in terms of levels and localization. A strain was considered to be a responder in terms of abundance if $KS_{\text{stat}} \geq 0.3$ for at least two of the three replicates and $f \geq 1.5$ (for induced proteins) or $f \leq 0.75$ (for repressed proteins). $KS_{\text{stat}} \geq 0.3$ for at least two of the three replicates was also used for NCR changes. For KS tests replicates were compared pairwise. For protein levels, no AF correction is performed it is not possible to determine AF for each cell. We also kept track of fold changes and subcellular localization as scored by Huh et al. (15) so that true translocations and NCR changes due to abundance changes can be distinguished.

The Cytoscape program was used for all network analyses (22,23). GO analyses were performed using the ClueGO plugin in Cytoscape (24). The YEASTRACT website was used for TF analyses (http://www.yeastract.com/) (25,26). Further details of statistical methods used can be found in the Supplementary Information File.

RESULTS
Sample preparation and analyses
We set out to monitor expression and localization changes for 4159 GFP-tagged proteins in the same number of $S$. cerevisiae strains after 3h of exposure to 0.02% MMS, a relatively non-toxic dose (4,8). In the parental strain, cells showed an expected S-phase arrest and were largely viable at this time point, although the culture showed a $\sim 40\%$ decrease in colony forming ability (Supplementary Figure S8–S10). To address a specific timepoint across large numbers of samples using multi-spectral fluorescence imaging of distinct cellular compartments, we developed a fixation strategy to minimize the typical attenuation of GFP signal resulting from standard yeast fixation protocols. This was particularly important to detect changes in expression levels for low copy-number proteins that are otherwise obscured by cellular autofluorescence. In a previous study of the 4159 strains expressing the GFP-tagged proteins, only 2700 proteins could be reliably detected using live-cell flow cytometry (16), and of these, the vast majority (85%) of GFP levels resided at the low-end of the expression range (16) (Supplementary Figure S1). We therefore developed an optimized aldehyde-based fixation protocol that minimized intensity loss and, importantly, exhibited a linear relation between fixed and live cell intensities across the entire range of GFP expression so that relative changes upon MMS exposure could be accurately measured (Supplementary Figures S2–S7). Strains were grown in minimal media in triplicate using a robotic liquid handler (see ‘Materials and Methods’ section for details). The importance of replicate measurements for generating greater statistical confidence in results from high-throughput screens has previously been emphasized (27), especially for detecting small but biologically significant responses.

Early log-phase cultures were incubated in control medium or medium containing 0.02% MMS for 3h. Cell were then fixed, and nuclei and cell-walls were stained. (Figure 1A). Custom-written image analysis programs were used to quantify GFP fluorescence in segmented cytoplasmic and nuclear compartments (Figure 1B, ‘Materials and Methods’ section). The computed intensities from fixed cells in this work compared well with previous studies on live cells (Supplementary Figures S11 and S12). Finally, statistical analyses were performed to identify ‘responders’ defined as strains for which protein levels were significantly induced or repressed, as well as strains for which proteins were enriched in the nucleus or cytoplasm as quantified by their NCR. Significant differences in mean expression or NCR were determined using a Student’s t-test with significance value of $P < 0.05$ (Figure 1C). However, because the comparison of mean values alone does not reveal differences in the underlying protein distributions themselves, the KS test was also employed (28–31). The KS statistic has been shown to be significantly more sensitive in detecting differences in population responses compared with the comparison of mean responses (29). The KS statistic represents the maximum distance between two cumulative histograms, where values equal to or greater than 0.2 are suggested to denote biological significance in the context of HCS (28,29). In addition to $t$-tests comparing mean responses over replicates, we used the KS statistic to identify responders in terms of abundance or NCR changes (Figure 1C and D; ‘Materials and Methods’ section). ‘Hit-lists’ for induced and repressed proteins, as well as nucleus- or cytoplasm-enriched proteins, are provided in Supplementary Table S1 as measured by both t-tests and KS tests, with a summary of some major responders presented in Table 1. Also listed
is the subcellular localization of each protein, as previously determined (15), and its function as listed in the Saccharomyces Genome Database. Overall, 415 induced proteins, 174 repressed proteins, 133 nuclear-enriched proteins and 10 cytoplasm-enriched proteins were identified.

Supplementary Table S2 documents intensity and NCR measurements for every strain measured.

Induced proteins and their interaction networks represent a wide range of cellular processes including DNA repair, proteolysis, chromatin remodeling and ribosome biogenesis.

GO functional enrichment analysis for the 415 induced proteins using the ClueGO plugin in Cytoscape (22–24) revealed an expected enrichment of GO-terms related to DNA damage response, DNA repair and cellular stress, as well as other GO-terms related to cellular metabolism, protein degradation, chromatin remodeling, RNA polymerase II transcription, mRNA processing and ribosome biogenesis (Supplementary Table S3 and Figure 2A). Although the absolute number of genes associated with some GO terms can be low, they may still be scored as significant when they represent a large fraction of the total genes associated with that term. For example, 8/14 (57%) proteins associated with the ubiquitin-independent proteasomal machinery, and 5/7 (71%) proteins associated with trehalose metabolism are induced by MMS treatment (Figure 2A). The first indication that proteasome function is enhanced and that trehalose metabolism is affected upon MMS exposure came from transcriptional profiling studies (6,7). Subsequent work has now shown that proteasome-mediated responses are directly involved in DNA repair (32), and that trehalose protects cells against different DNA damaging agents (33–35). Thus, although these and the other GO-enriched processes may initially seem disparate, taken together they represent a coordinated response to cellular insult by MMS.

Mapping responders onto a previously compiled yeast interactome (36) enabled identification of functional networks of induced proteins. The full interactome annotates physical, genetic and TF interactions. The smaller network that resulted from mapping MMS-induced proteins onto the full interactome had higher connectivity, higher clustering coefficient, and a greater ‘Large Connected Component’ (LCC) than random networks of
The top 20 responders (where applicable) in the induced, repressed, nucleus- and cytoplasm-enriched categories are shown. In the blue rows are significant responders by t-tests, and in the pink rows are the significant responders by KS tests. For the induced and repressed categories, the responders are arranged according to maximal fold change. For the nucleus- or cytoplasm-enriched categories, responders are arranged by maximal NCR change for t-tests and by maximal KS statistic for KS tests. For the complete list of responders, see Supplementary Table S1.

similar features were previously observed in network analyses of toxicity modulating genes in genomic phenotyping studies (37). Although the protein–protein and genetic interaction maps revealed a number of clusters that were expected from the GO functional enrichment analysis (namely, DNA damage response, ribosome biogenesis, chromatin remodeling, RNAPII transcription and proteolysis), such mapping also highlighted additional functions that were not represented in the GO analysis (Figure 2B). For example, all four proteins known to be associated with the cohesin complex (38) are induced. This group was not represented in the GO analysis because the number of proteins in the cohesin complex is so small that it falls below the threshold used for filtering GO terms. It turns out that previous studies also demonstrated a role for the cohesin complex in DNA damage response beyond its conventional role in sister-chromatid cohesion (39–41). We also identified a distinct cluster of heat shock proteins (Figure 2B), consistent with recent studies that demonstrate links between heat shock and DNA damage responses (42,43). Finally, a group of proteins involved in mRNA processing was present among the subnetworks of induced proteins. Interestingly, a recent genomic phenotyping study involving both essential and non-essential genes also highlighted the importance of mRNA processing and splicing proteins in governing sensitivity to the toxic effects of MMS (8). MMS also induces damage to proteins, RNA and other cellular components, in addition to DNA. The modulation of a large number of RNA processing and ribosomal genes may also be signaled from protein and RNA damage in conjunction with the direct DNA damage response.

**Partitioning of biological function among induced proteins according to number of upstream TFs**

In addition to physical and genetic interactions, we examined TF interactions by mapping induced proteins onto the global TF network constructed from the YEASTRACT database (25,26,36), thereby identifying eight TFs that are themselves induced (Yap1, Cst6, Cin5, Dot6, Xbp1, Pho2, Dal81—see red text in Figure 3A). These eight TFs collectively have the potential to regulate the expression of 54% of the induced proteins (Figure 3A). Deletion mutants for five of the eight TFs (Yap1, Cst6, Cin5, Dot6, Dal81—circled in green in Figure 3A) showed MMS sensitivity in the genomic phenotyping assay (5), and Xbp1 is known to be a stress-responsive TF. In general, 11% of all possible targets for these eight TFs are represented among the induced proteins. Although this is higher than the 6.5% that would be expected for random networks of the same size, it is still a relatively small fraction of all possible targets. Although this may be due in part to the fact that the GFP library represents only 70% of the yeast genome, it more likely indicates a tendency for cells to require combinations of TFs to modulate gene expression in response to environmental challenges, rather than allowing promiscuous non-specific upregulation of all possible targets for a single TF. Although a number of the induced proteins are targeted by more than one of the eight TFs (i.e. have an indegree >1, Figure 3A), these eight MMS-induced TFs by no means represent all of the TFs that are potentially capable of regulating the induced proteins.

To obtain a comprehensive picture of all TFs that might govern the expression of induced proteins, irrespective of whether the TFs themselves were induced, we used the updated YEASTRACT database that documents all known major TF interactions in the yeast genome (25,26). This analysis identified 59 TFs, each of which potentially governs the expression of at least 5% of the induced proteins. Most induced genes were governed by more than one of the 59 TFs, with four as the median value of node in-degree, which represents the number of TFs governing the expression of a target protein (Figure 3B).
To test whether the induced protein nodes that can be governed by a large number of TFs serve distinct biological functions compared with more isolated nodes, we filtered out genes that can be targeted by four or fewer of the 59 putative TFs (representing proteins that are regulated by relatively few TFs) from genes that can be targeted by more than four TFs (representing proteins that are regulated by many TFs). GO analysis for these two sets of MMS-induced proteins revealed a clear difference in functional enrichment, with \( P \)-values as low as or lower than those for the combined set of all induced proteins shown in Figure 2A (Figure 3C; Supplementary Table S4). Interestingly, induced proteins associated primarily with metabolic processes were found to be targeted by five or more TFs, whereas DNA damage response and chromatin remodeling proteins represent the more isolated nodes that are targeted by four or fewer TFs. In the full transcription network, the median value for indegree is also four, which is similar to the subset of MMS-induced proteins (Supplementary Figure S14). Not surprisingly, when genes are partitioned as mentioned previously in the full network, DNA-related processes do not dominate the more isolated nodes because all cellular functions are now represented. However, metabolic processes continue to be over-represented among the nodes with five or more upstream TFs, indicating that this may be a generic feature of the transcriptional network (Supplementary Figure S14).

**Enrichment of lipid biosynthesis and membrane trafficking processes among repressed proteins and networks**

In addition to induced genes, transcriptional profiling studies have revealed many genes whose transcripts are downregulated by MMS (6,7,9). At the protein level, one might expect fewer repression responders due to the longer half-lives of proteins compared with mRNAs, except in the case of targeted protein degradation. In other words, even for a gene that is transcriptionally silenced, its protein products may remain in the cell for some time, particularly under conditions of inhibited growth. In our screen, 174 repressed proteins were identified, even with a lower cut-off threshold of 25% reduction compared with the 415-induced proteins that showed >50% induction. \( P \)-values for GO-function enrichment were generally larger for repressed versus induced proteins (Supplementary Table S3 and...
Figure 4A), and the network of repressed proteins had lower connectivity than the induced proteins as measured by their connectivity that was comparable with random networks of the same size (Supplementary Figure S15). However, cellular membrane organization, secretion, trafficking and peroxisome organization were significantly over-represented among the repressed proteins (Figure 4A and B). Also, the single repressed TF, Phd1, is known to control a number of sterol biosynthesis genes known to be important for lipid and membrane biosynthesis (Figure 4C). Finally, several proteins involved in chromatin remodeling and regulation of the mitotic cell-cycle were also part of the network of repressed proteins (Figure 4B), even though none of these processes was significantly enriched in the GO analysis (Figure 4A).

We used the YEASTRACT database to identify 52 TFs that can target the expression of 5% or more of the repressed proteins. Most of these (47/52) are found among the TFs that target genes for the induced proteins, presumably because some of these TFs (e.g. Yap1, Sfp1 and Ste12) have high numbers of targets in the genome (Figure 4D). However, although we observed highly significant enrichment of the targets of these TFs for the induced proteins ($P = 1.8 \times 10^{-15}$ overall, Figure 4G), the enrichment was not nearly so high ($P = 9.4 \times 10^{-4}$ overall, Figure 4G) for repressed proteins (Figure 4E–G). The number of targets for each TF in both lists was divided by the number of targets expected by random chance from the whole genome, to assess specific enrichment of the targets for a given TF in each list. This measures the normalized occurrence for each TF (thus a normalized occurrence of 1.5 indicates that 50% more targets are present in the induced or repressed list when compared with random lists of the same size that exhibit a mean normalized occurrence of 1).
analysis revealed the targets of TFs for all three major stress-responsive pathways in yeast [genes with heat shock elements, stress response elements or AP-1 responsive elements (6,9,44–46)] to be particularly enriched among the induced proteins. TFs Yap1, Msn2, Msn4, Rpn4, Hsf1, Met4 all exhibit a high normalized occurrence among the induced proteins (Figure 4E, dotted circle). There were other TFs that showed yet higher normalized occurrence, but the absolute number of targets for these TFs in the genome was low, and in effect, they targeted only several of the induced genes.

Signatures of translational regulation of induced proteins

Cells use both transcription and translation to regulate gene expression. Transcriptional responses to DNA damaging agents are well documented (6,9), and recent studies have unveiled a translational component to such responses (12,13). The study by Begley et al. (2007) computationally identified a set of 425 genes with a skewed codon usage pattern such that their translation would be promoted by the Trm9 tRNA methyltransferase that catalyzes specific tRNA modifications that change codon-anti-codon affinity. Such modifications affect the efficiency of translation for a subset of transcripts rich in specific codons, especially under conditions of DNA damage (13). Ribosomal, metabolism and stress response genes were enriched in this group of 425 potential preferentially translated (PPT) genes. However, belonging to the PPT group does not ensure induction under conditions of damage, for translational components can offset translational responses. Despite this, we find 57 PPT proteins...
among the MMS-induced proteins, approximately twice the number that is expected by random chance (Supplementary Figure S16A). These 57 proteins form a close network with several RP genes; GO analysis reveals enrichment of processes related to carbohydrate and trehalose metabolism, ribosome biogenesis, oxidative stress and deoxyribonucleotide production (Supplementary Figure S16B and C).

Nucleus and cytoplasm enriched proteins and networks

The list of proteins that become enriched in the nucleus in response to MMS includes two categories: (i) nuclear proteins whose relative expression increases in response to MMS and (ii) proteins that translocate from the cytoplasm to the nucleus. Although not all induced nuclear proteins are represented in this list, instances where the NCR increases simply reflects the induction of a protein were retained because they captured subtle expression increases in the nucleus not represented in the total list of induced proteins (Supplementary Table S1). For example, Ubi4, yeast ubiquitin, is scored as both an induction and a nuclear enrichment responder, but Rxt2, a subunit of the histone deacetylase Rpd3 complex implicated in the activation of DNA damage induced genes (47) is found only in the nuclear enrichment list. Remarkably, GO analysis of this relatively small list of proteins (133) produced extraordinarily high-significance values for functional enrichments, and perhaps not surprisingly, most of the nuclear-enriched functional categories pertained to DNA and RNAPII-related processes, with a notable absence of the metabolic, proteasomal and ribosomal processes seen in the total list of inductions. P-values for functional enrichment (Supplementary Table S3 and Figure 5A) were substantially lower even than those for the induced proteins as a whole (Figure 2A), and the network of nuclear-enriched proteins exhibited connectivity far greater than what would be expected for similarly sized random networks, reflecting the close functional interactions among these proteins (Supplementary Figure S17). For several GO terms associated with the nuclear-enriched proteins, >25% of all associated proteins were represented (Figure 5A). The physical and genetic protein interaction networks were dominated by components of chromatin remodeling, RNAPII-dependent transcription, plus mRNA and snoRNA processing proteins. Components of the cohesin complex were also present among the nuclear-enriched proteins.

TF network analysis identified three nuclear-enriched TFs (Ste12, Abf1, Dot6) that regulate a number of proteins in the list of nuclear-enriched proteins. Although other TFs such as Ixr1 previously implicated in the DNA damage response (48,49) were not part of this network, they were present in the list of nuclear enrichments (Supplementary Table S1). These are instances of subtle inductions that were only seen in the nucleus but missed in the total list of induced proteins. Conversely, two proteins that are a part of the induction list and known to be nuclear-enriched upon damage (Ubc13, Yap1) were not found in this study. Although Yap1 with a KS-statistic value of 0.28 lay just below the cutoff of 0.3 for nuclear enrichment, it should be noted that in previous work, more than a 10-fold higher MMS dose was used to induce its nuclear translocation, compared with the present study (50). Thus, nuclear enrichment of a protein can be dose-dependent.

Evidence from several distinct lines of work have demonstrated connections between chromatin remodeling and the DNA damage response in eukaryotes (4,6,7,9,47,51–53). Our data (Figures 2B and 5B) also indicate extensive changes in proteins for the remodeling of chromatin and for nucleosome disassembly that are mobilized on exposure to MMS, presumably to allow repair machinery access to sites of DNA lesions.

Surprisingly, few cytoplasmic enrichments (10 proteins) were identified in our screen (Supplementary Table S1). This may in part be due to the fact that the cytoplasmic volume is significantly larger than the nuclear volume, rendering it difficult to detect translocation of low-expressed proteins. However, even amongst this small number of proteins, interesting features stand out. For example, we find that the Wtm1 and Wtm2 proteins translocate out of the nucleus into the cytoplasm in response to MMS, as does Rnr4 (Figure 5D and E). Wtm proteins are involved in nuclear anchoring of the RNR small-subunits (54–57), with one study previously implicating the Wtm proteins in the control of RNR transcription (58). It appears that controlling the subcellular localization of the Wtm proteins may provide an additional mode of RNR regulation.

RP response to MMS is dose dependent

A surprising GO category for induced proteins was ribosome biogenesis and components of the ribosomal machinery because previous studies from several groups have shown that ribosomal genes are generally transcriptionally repressed under conditions of DNA damage (6,7,9,10). The regulation of ribosomal genes is thought to be primarily at the level of transcription in yeast (59), but almost 90% of all RP genes are also found in the list of PPT genes (13), suggesting cells use both transcription and translation to tune ribosome numbers (also see Supplementary Figure S16). A recent study similar to ours (discussed in more detail later in the text) also identified this group among induced proteins (20). To investigate this apparent discrepancy, we examined two such induced RPs (Rpl7a-GFP and Rps22a-GFP) using live-cell flow cytometry. We found that at moderate doses of MMS (0.02%, 2 h), both proteins were induced, whereas at higher MMS dose (0.1%, 2 h), there was a small but significant repression of the proteins (Figure 6A). The 2-h time-point was chosen because it was intermediate between the 3-h time-point at which the present screen is performed, and the 1-h time-point of the first transcriptional profiling study that revealed transcriptional repression of the ribosomal genes at 0.1% MMS (7). This suggests, not surprisingly, that cellular response differs according to the extent of damage, presumably depending on whether the cell can repair damage and proceed through the cell-cycle, or whether growth halts completely. A similar pattern of induction was seen in
Tandem Affinity Purification (TAP)-tagged strains of Rpl7a and Rps22a (Supplementary Figure S18). Because ribosome biogenesis is intimately linked with cell growth, we investigated how cell growth characteristics differ at the two MMS doses.

Using both liquid and agar assays, we found that at 0.02%, MMS cells proceed through the cell-cycle, albeit slowly, whereas at 0.1%, MMS cell growth is abrogated entirely (Figure 6B and C, Supplementary Figure S19). Conditions of general cellular stress inhibit growth, and at the same time, ribosome biogenesis in a Target of Rapamycin (TOR)-dependent manner (17,59). The TOR pathway has also been shown to control DNA damage responses by controlling dNTP production (60), and the TOR pathway effectors Sch9 and Sfp1 are known to be involved in both ribosome biogenesis and stress responses (17,61,62). Sfp1 is a nuclear-localized TF that regulates RP expression and translocates to the cytoplasm in response to various stresses including DNA damage by MMS (0.1%), thus turning off RP gene expression (17). Sfp1 itself is also induced by MMS damage, which may appear to conflict with the aim of shutting down ribosome biogenesis (63). We therefore investigated the levels and localization of Sfp1-GFP in response to moderate and high doses of MMS. At both doses, Sfp1-GFP was induced as assessed by live-cell flow cytometry (Figure 6D). Indeed, although Sfp1-GFP showed only a 27% increase in the original screen, and was not scored as induced (only proteins with >50% increase in expression were considered to be responders), this small increase was significant ($P = 0.003$). Expression of Sfp1-GFP in the absence of damage is low, making estimations of fold change difficult. Our conservative estimates of fold change in the global screen are systematically underestimated when the fluorescence signal is close to cellular autofluorescence (see ‘Materials and Methods’ section), causing Sfp1-GFP to be absent from the list of induced proteins. However, when observed with higher resolution microscopy, at the moderate MMS dose (0.02%), Sfp1-GFP was induced and clearly nuclear, thus being available to upregulate the expression of RP genes. At the higher dose (0.1% MMS), Sfp1-GFP, although still present at induced levels, became cytoplasmic, concomitant with the repression of ribosomal genes. This
provides a clear example where tuning both the abundance and the nuclear-versus-cytoplasmic localization of a protein can be used by cells to effect induction and repression of gene expression.

**DISCUSSION**

**Identification of repressed proteins**

The present study provides a comprehensive single-cell-level view of an orchestrated cellular response to damage induced by MMS. Previous studies using the yeast GFP fusion library to interrogate protein level changes in response to MMS, by imaging (20) or flow cytometry (21) used a single replicate of live cells. Here, fixation of cells was essential to ensure a specific and consistent exposure time to the damaging agent, as well as unambiguous identification of the cell nucleus and cell-wall without expression of an additional fluorescent protein as a nuclear marker. In previous studies (20,21), no proteins met the cutoff set for downregulation, even though many genes are known to be transcriptionally repressed in response to MMS (6,7,9). As discussed, protein level repressions may manifest as small changes at early time points after damage exposure, and hence replicate measurements may be necessary to identify them reliably. Similar doses and time points were used in all these studies: Lee et al. used 0.02% MMS for 4 h; Tkach et al. used 0.03% MMS for 2 h, whereas we use 0.02% MMS for 3 h. Here, triplicate measurements enabled the identification of a number of proteins whose expressions are reduced on DNA damage. Processes of membrane-trafficking, lipid synthesis and peroxisome function were enriched among the repressed proteins. In breast cancer models, mutations in the critical tumor suppressor protein p53 have been associated with expression of sterol biosynthesis genes (64). In mouse liver, activation of the protein Nrf2 by oxidative stress has been shown to be associated with downregulation of lipid biosynthesis, to allow the scavenging of reactive oxygen species (ROS) by reduced nicotinamide adenine dinucleotide phosphate (NADPH) that functions both in lipid metabolism pathways and in

**Figure 6.** Differential response of RPs to different levels of MMS damage. The response of two candidate induced RPs (Rpl7a-GFP, Rps22a-GFP) to MMS damage was investigated with live-cell flow cytometry, at 2 h. (A) At 0.02% MMS, the RPs are induced, whereas at the higher dose (0.1%), they are repressed. (B) Parental BY4741 cells were subjected to similar treatments, and culture densities (events/ml) were determined by flow cytometry (optical density measurements are expected to be not accurate at these early time points, as MMS arrest can cause an increase in cell-size). For three early log-phase cultures, initial densities were measured, and a third of each culture was subjected to 0, 0.02%, 0.1% MMS treatment. All densities at 2 h are normalized by the density of the initial cultures. 0.02% MMS significantly retards growth, but does not abrogate it altogether, whereas there is almost no growth in the presence of 0.1% MMS. (C) In a different assay from liquid cultures, even for cells growing on agar-pads with or without MMS, there is slow growth with 0.02% MMS, whereas no growth is seen at 0.1% MMS. Images of the same fields at 0 and 15 h are shown. (D) The GFP-tagged TF Sfp1, which controls RP gene expression, is induced at both doses of MMS as determined by live-cell flow cytometry. AF-corrected intensities are plotted; however, the expression of the protein is low in the absence of damage, making estimations of fold change difficult. (E) At 0.02% MMS, the protein is still nuclear, whereas at 0.1% MMS, it is clearly cytoplasmic, concomitant with the induction and repression of the RP proteins. Live cells were imaged with a 100× oil-immersion objective, given the low expression of the protein. Overlays of GFP and phase images are presented. The scalebar is 5 μm.
ROS scavenging (65,66). Peroxisomes may act as a source of ROS that affect nucleic acids, proteins and lipid, and they are kept under tight control in yeast (67); indeed, peroxisome proliferation causes oxidative DNA damage in rat livers and plays a role in hepatocarcinogenesis (68). Although these findings span a number of distinct model systems, they provide plausible explanations for the genes repressed by MMS treatment, as seen in this study, and indicate that these may be general features of regulation under conditions of damage.

Comparisons with other studies

Despite the attenuation of GFP signals on fixation, the intensities measured in fixed cells in this work compared well with previous studies on live cells performed with both flow cytometry and imaging methods (16,20) (Supplementary Figures S11 and S12). Further, MMS-induced protein groups are largely similar between this study and that by Tkach et al. (20). In terms of protein localization, manual annotation of protein localization in the prior study allowed finer discrimination of subcellular localization into organelles, but at the cost of replicate measurements. In comparison, here, we measure broad nuclear versus cytoplasmic localization in a fully automated manner and identify several highly nuclear-enriched DNA-related processes, as well as abundance changes that span a wide variety of cellular responses. Although we lose detail in the granularity of subcellular organelles, we detect GFP intensity automatically in the full nuclear and cytoplasmic masks, combined with greater statistical confidence from replicate measurements.

The number of responders identified in this study is substantially fewer than the transcriptional profiling studies. However, comparison with an early transcriptional profiling study from our laboratory (7) reveals that 62% of all the induced proteins are transcriptionally induced in response to MMS. In contrast, only 6.3% of the repressed proteins were found to be transcriptionally repressed. Indeed, although correlations between transcript and protein levels in unperturbed cell populations are poor (69,70), it has recently been shown that under conditions of environmental stress, transcript induction correlates well with protein induction, but transcript reduction produces negligible change in protein levels (71). Protein level repression under conditions of DNA damage presumably occurs due to targeted degradation (72) instead of transcriptional repression. Several studies from our laboratory, including the present one, have indicated a key link between the proteasomal machinery and response to MMS-induced damage (6,73).

TF network analysis

Extensive TF analyses identified Yap1 as a major induced TF. Yap1 has many targets in the yeast genome, and its role in stress responses is well-established (45,50,74). In addition to causing direct alkylolation damage, MMS can also elicit an oxidative stress response in cells, and Yap1 was recently shown to be a major regulator of this response (74). We also found specific enrichment of the targets of several other major stress-responsive TFs (Rpn4, Msn2, Msn4, Hsf1 and Met4) among the induced proteins. Combinations of a large number of TFs were upstream of the observed metabolic response genes, whereas fewer TFs directly targeted genes for DNA-related processes.

It should be borne in mind that the network analyses performed here projects responders onto static protein–protein, genetic and TF interaction networks. It is thought that genetic interactions may shift on DNA damage, whereas physical interactions remain largely unchanged (75). TF combinations affecting a gene may also shift on DNA damage (49), and the periodically updated YEASTRACT database documents most known TF interactions under various conditions from different sources, based on both bioinformatic and experimental data. However, under any one condition, only a subset of the targets is likely to be active for a given TF.

DNA-related processes in nuclear-enriched proteins

Interestingly, nuclear-enriched proteins are found to form a closely connected network, with a corresponding significant enrichment of DNA-related processes that include transcription, chromatin remodeling and DNA repair, and a notable absence of proteins of the metabolic, ribosomal and proteolytic machinery. Thus, the list of nuclear-enriched proteins clearly segregates the central DNA level responses from the other cellular processes that together orchestrate the total cellular response to MMS insult.

Differential response of RPs according to MMS dose

Finally, differing responses for several RPs were observed depending on the MMS dose. These responses correlated with altered cell growth and the cytoplasmic translocation of the Sfp1 TF that results in shutting off RP gene expression. Although Sfp1 is a central player for RP gene expression, other factors also affect decisions to either stall growth or to repair damage and proceed through the cell cycle. Recently, proteins of the cohesin complex have been shown to be involved in RP gene regulation (76), in addition to direct DNA damage responses (39,40). The cohesin complex proteins are upregulated under the conditions of damage used here, which could in turn feed back onto RP expression. Future work will explore whether the upregulation of specific RPs is for the replacement of damaged ribosomal components (77,78) or to make functionally specialized stress-specific ribosomes (79,80) or perhaps both.

Outlook

Taken together, our work presents a global systems-level proteomic view of the cellular response to MMS damage. Transcription may represent the first level of regulation, but now, we reveal the protein responders in terms of levels and broad subcellular localization. Although focused studies investigate a few genes in isolation, intricate connections within the yeast proteome suggest that no response can be isolated from cascading effects within the network. Yet, despite the seemingly daunting complexity, the overall picture that emerges reveals a coordinated response in terms of DNA repair, chromatin remodeling,
proteolysis and cell growth, indicating that the system is tuned to buffer a fairly large range of genotoxic challenges.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online, including [81–95].

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