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Comprehensive Characterization of Genes Required for Protein Folding in the Endoplasmic Reticulum

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Protein folding in the endoplasmic reticulum is a complex process whose malfunction is implicated in disease and aging. By using the cell's endogenous sensor (the unfolded protein response), we identified several hundred yeast genes with roles in endoplasmic reticulum folding and systematically characterized their functional interdependencies by measuring unfolded protein response levels in double mutants. This strategy revealed multiple conserved factors critical for endoplasmic reticulum folding, including an intimate dependence on the later secretory pathway, a previously uncharacterized six-protein transmembrane complex, and a co-chaperone complex that delivers tail-anchored proteins to their membrane insertion machinery. The use of a quantitative reporter in a comprehensive screen followed by systematic analysis of genetic dependencies should be broadly applicable to functional dissection of complex cellular processes from yeast to human.

The endoplasmic reticulum (ER) is responsible for the folding and maturation of secreted and membrane proteins. External stress or mutations can compromise ER folding, contributing to diseases such as diabetes and neurodegeneration (1, 2). The specialized milieu of the ER is composed of a large number of proteins that aid the structural maturation of itinerant proteins (3, 4). Although many of these ER folding factors

have been extensively studied, the full range of proteins contributing to this process is unknown, and how they function together is poorly understood.

Systematic identification of genes contributing to ER folding. We exploited the cell's endogenous sensor of ER protein folding status, Ire1p, to identify genes in *Saccharomyces cerevisiae* that contribute to structural maturation of secretory proteins. In response to misfolded proteins in

the ER, the transmembrane sensor Ire1p activates the transcription factor Hac1p (5), which in turn transcriptionally up-regulates a distinct set of genes (6, 7) in a process called the unfolded protein response (UPR). We used a reporter system in which a Hac1p-responsive promoter drives green fluorescent protein (GFP) expression (8) (Fig. 1A). To correct for nonspecific expression changes, we coexpressed a red fluorescent protein (RFP) from a constitutive *TEF2* promoter and used the ratio of GFP/RFP as our reporter of UPR signaling. A titration of the ER stress-inducing reducing agent dithiothreitol (DTT) demonstrated that this reporter quantitatively responds to misfolding of ER proteins (Fig. 1B).

With use of synthetic genetic array methodology (9), we introduced the reporter into ~4500 strains from the *S. cerevisiae* deletion library (10),

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whose deletion induces the UPR and the genes that were previously shown by microarray analysis to be transcriptionally up-regulated by the UPR (7) [fig. S2 and table S2; see also (10, 13)]. Thus, although defining the UPR targets was fundamental to our understanding of how cells respond to ER stress, it provides a limited view of the processes constitutively required for folding in the ER.

Overview of gene deletions affecting the UPR.

Proteins whose deletion caused up-regulation of the reporter were highly enriched for localization throughout the secretory pathway (fig. S3 and table S3) (14), including the ER as well as the Golgi, vacuole, and endosome. As expected, chaperones (15) and genes in the N-linked glycosylation (16) and ER-associated degradation (ERAD) (17) pathways featured among the top hits (Fig. 1C). However, deletion of genes involved in many other processes, including O-mannosylation, glycoposphatidylinositol anchor synthesis, lipid biosynthesis, multiple steps of vesicular trafficking, and ion homeostasis, caused similarly high reporter inductions (table S1). Moreover, the UPR up-regulators included several dozen poorly characterized genes, some whose deletion caused reporter induction that rivaled the strongest hits.

The diversity of functions contributing to ER integrity presented a major obstacle in our efforts to understand how these unexpected factors func-

tion together to support protein folding in the ER. To overcome this, we explored their functional dependencies by systematically quantifying how the phenotype caused by loss of one gene was modulated by the absence of a second. Systematic analysis of genetic interactions, using growth rate as a phenotype, has been used extensively to determine gene function (9, 18–23). We sought to generalize this strategy by using ER stress as a quantitative phenotype. Accordingly, we quantified GFP reporter levels in over 60,000 strains containing pairwise deletions among 340 of our hits (12).

Genetic interactions illuminate functional relations. Three examples illustrate the utility of the double mutant analysis. First, comparison of the GFP levels in the presence and absence of *IRE1/HAC1* differentiated between the subset of up-regulators whose deletion affected protein folding in the ER (i.e., GFP induction was dependent on the Ire1p/Hac1p pathway) from those, like chromatin architecture genes, that were directly affecting expression of the reporter (Fig. 2A and table S4).

In a second example, UPR levels of pairwise deletions with $\Delta die2/alg10$, the enzyme that performs the last step in the synthesis pathway for N-linked glycans, illustrated our ability to define genes acting in a linear pathway (Fig. 2B). Most double mutants showed a typical increase

in fluorescence that was dictated solely by the reporter levels of the single mutants. Notably, a specific subset of the double mutants had the same reporter level as the single mutant, indicative of “fully masking” epistatic interactions in which the function of one gene is completely dependent on the presence of a second one. Indeed, the genes that we find to be epistatic to *DIE2* include the full set of factors that act immediately upstream of Die2p in the synthesis of N-linked sugars (16).

The utility of aggravating genetic interactions, in which pairs of deletions lead to exaggerated folding defects, was illustrated in a third example in which we overexpressed the constitutively misfolded and rapidly degraded membrane protein KWS (24) in deletion strains that were hits in our screen (Fig. 2C). KWS degradation is mediated by a well-defined subset of the ERAD machinery, including the E3 ubiquitin ligase Ssm4p/Doa10p and the E2 ubiquitin-conjugating complex Ubc7p and Cue1p (24). The role of these factors in mitigating the stress caused by overexpression of KWS is revealed in our data by the strongly aggravating interaction between their deletions and KWS overexpression. In contrast, other ERAD components, which do not act on KWS, including the Hrd1p/Hrd3p E3 ligase complex (24), show typical reporter levels.

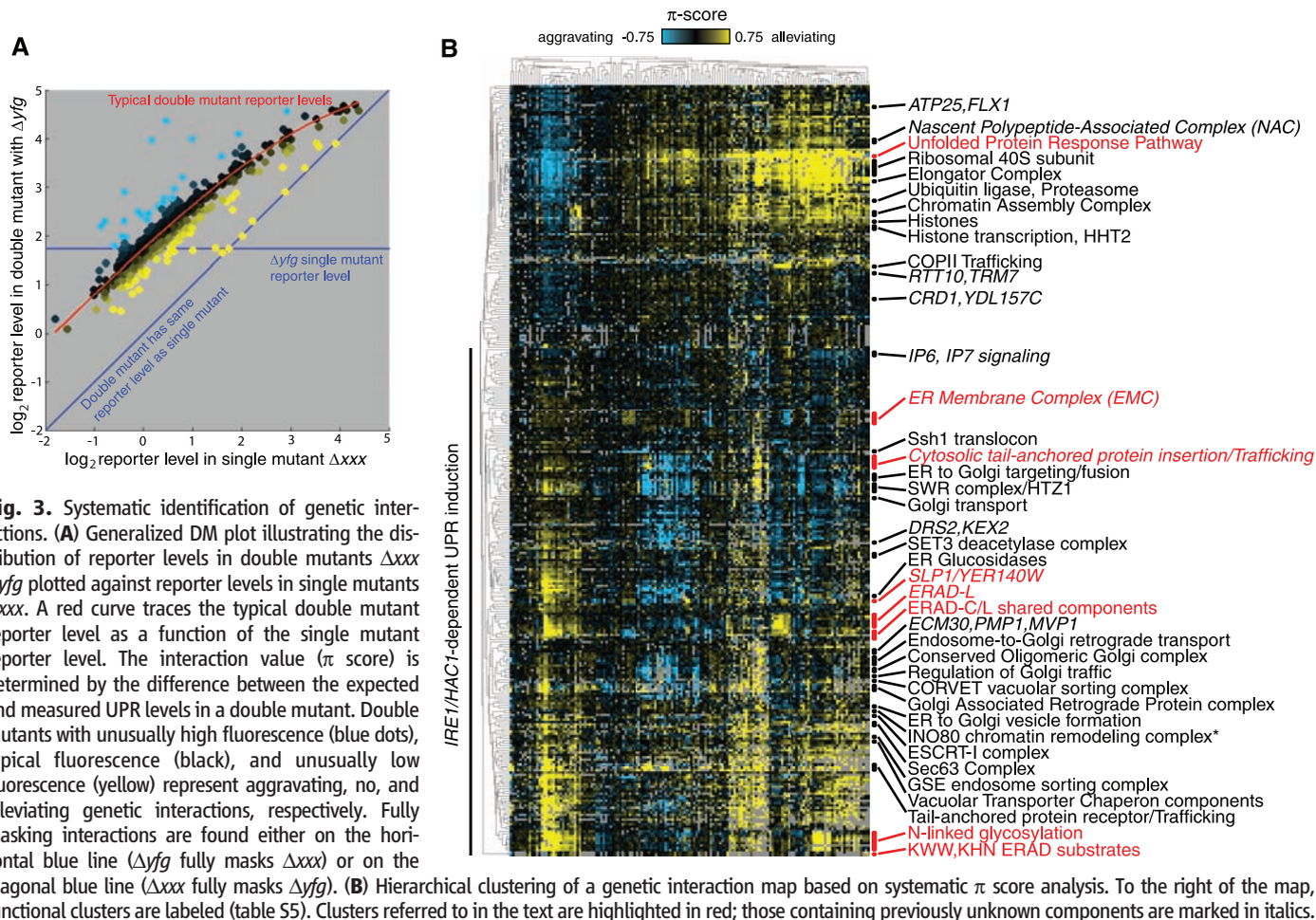


Fig. 3. Systematic identification of genetic interactions. **(A)** Generalized DM plot illustrating the distribution of reporter levels in double mutants $\Delta xxx \Delta yfg$ plotted against reporter levels in single mutants Δxxx . A red curve traces the typical double mutant reporter level as a function of the single mutant reporter level. The interaction value (π score) is determined by the difference between the expected and measured UPR levels in a double mutant. Double mutants with unusually high fluorescence (blue dots), typical fluorescence (black), and unusually low fluorescence (yellow) represent aggravating, no, and alleviating genetic interactions, respectively. Fully masking interactions are found either on the horizontal blue line (Δyfg fully masks Δxxx) or on the diagonal blue line (Δxxx fully masks Δyfg). **(B)** Hierarchical clustering of a genetic interaction map based on systematic π score analysis. To the right of the map, functional clusters are labeled (table S5). Clusters referred to in the text are highlighted in red; those containing previously unknown components are marked in italics.

Phenotypic interaction score (π score) quantifies functional relations. Strongly aggravating and fully masking interactions as described above are only a subset of the broader range of possible genetic interactions in which pairs of perturbations lead to a continuum of exacerbated (aggravating) or attenuated folding defects. We sought to quantify these systematically by developing a phenotypic interaction score or “ π score” that describes the degree to which a double mutant UPR reporter level differs from that expected from the two single mutant levels. A simple empirical multiplicative model accurately predicted the typical double mutant reporter levels when we accounted for saturation of the reporter [Fig. 3A; see (12)]. The π score for each double mutant was given by the difference between the typical levels expected from the reporter levels of the two single mutants and the measured UPR reporter levels in the double mutant. Thus, negative π scores (exaggerated inductions) represent aggravating interactions, and positive π scores (unusually low inductions) represent alleviating interactions, with the fully masking interactions being a subset of the positive π scores.

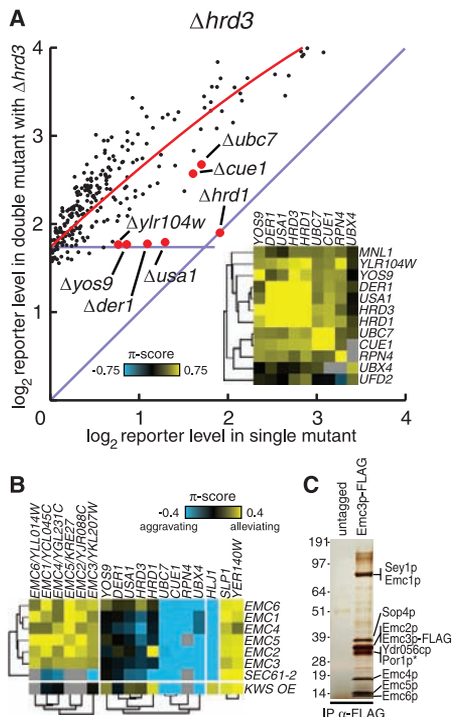
Systematic identification of functional groups through phenotypic interaction maps. In growth-based studies, the pattern of genetic interactions of a mutation provides a signature that can be used to

group genes by function (20, 21, 25). Analogous hierarchical clustering on the double mutant π scores yielded a map with a high density of precise functional clusters (Fig. 3B and fig. S4). This analysis accurately grouped over 100 of the previously well-characterized genes into 22 functions spanning a wide range of processes (table S5). Among genes whose deletions directly affected the ER folding environment (i.e., caused Ire1p/Hac1p-dependent reporter induction), our map grouped not only the ERAD and glycosylation machinery discussed above but also many other processes, including those in the distal secretory pathway (Fig. 3B). Our map also accurately clustered multiple functions that act downstream of *HAC1*, including the chromatin assembly complex, core histones, and histone chaperones.

Genetic interactions identify functional hierarchies. Within the functional groups defined above, the specific double mutant phenotypes revealed the extent to which the activities of individual components depended on each other. For example, all of the known components of the ERAD-L machinery needed for disposal of misfolded luminal proteins (17) formed a tight cluster. The double mutant phenotypes of $\Delta hrd3$ revealed the expected full dependence of *YOS9*, *DER1*, and *USA1* on *HRD3*

(Fig. 4A) (17). In contrast, only partial epistasis was seen with the E2 Ubc7p and its membrane anchor, Cue1p, consistent with their known roles in other branches of ERAD and the ability of another E2, Ubc1p, to partially substitute in their absence (26). In addition, the clustering analysis suggested that *YLR104W* is a previously unknown component of ERAD that acts upstream of *HRD3* and *USA1* (perhaps by delivering a subset of ERAD targets to the Hrd1p ligase) (table S6). Our complete list of genetic interactions, which includes over 500 full masking relations among 213 genes, should provide a resource of functional predictions for the community (table S6). For example, our data suggest that *YDR161W* is closely functionally related to the nascent polypeptide-associated complex. We also provide a MATLAB script to display double mutant plots for any gene in our data set (12).

Analysis of phenotypic interactions reveals previously unknown pathways important for ER protein folding. By using this systematic approach, we identified a pathway involving a conserved (table S9) multiprotein transmembrane complex. The poorly characterized genes *YCL045C*, *YJR088C*, *YKL207W*, *YGL231C*, *KRE27*, and *YLL014W* all clustered together and showed strongly alleviating interactions among themselves (Fig.



4B), a signature of factors that cooperate to carry out a single function (21). Immunoprecipitation of FLAG-tagged Ykl207wp revealed that proteins encoded by these genes form an apparently stoichiometric complex (Fig. 4C). Accordingly we termed this the ER membrane protein complex (EMC) and named the genes from this cluster *EMC1* through *EMC6*. Although the precise biochemical roles of the EMC will have to wait for future studies, our data suggest that loss of the EMC leads to accumulation of misfolded membrane proteins: The pattern of genetic interactions of strains deleted for EMC members most closely resembled that seen in a strain overexpressing the misfolded transmembrane protein Sec61-2p (a mutated form of the Sec61 translocon) (27) and is similar to the pattern of a strain overexpressing the misfolded transmembrane protein KWS (24). This shared pattern includes strong aggravating interactions with *Dubc7* and *Dcue1*, whose gene products are known to be involved in elimination of misfolded membrane proteins (17), but minimal interactions with other ERAD components.

A second cluster containing two conserved yet uncharacterized proteins (Yer140wp and Slp1p) show robust alleviating interactions with EMC components (Fig. 4B), as well as with each other. In support of a functional link between Yer140wp and Slp1p, these two proteins are suggested to be in a physical complex (28). The finding of two conserved protein complexes that are functionally dependent on each other underscores the value of this genetic data in identifying uncharacterized pathways required for ER folding.

Genetic interactions identify components of the tail-anchored protein biogenesis machinery. As a final example, we focused on Yor164c/Get4p and Mdy2/Tma24/Get5p because our analysis implicated them in tail-anchored (TA) protein biogenesis. TA proteins are an important class of transmembrane proteins, which includes soluble *N*-ethylmaleimide-sensitive factor attachment protein receptor (SNARE) trafficking factors (29, 30). TA proteins have a single C-terminal transmembrane domain, which is inserted into the ER membrane through the action of the recently discovered GET pathway: the Get3p/Arr4p adenosine triphosphatase (and its mammalian homolog Asna1/TRC40) binds newly synthesized TA proteins and brings them to the ER via the ER membrane receptor complex formed by Get1p/Mdm39p and Get2p/Hur2p/Rmd7p (31–33). Our double mutant analysis pointed to a role of Yor164c/Get4p and its physical interaction partner, Mdy2p/Get5p (34), in the GET pathway because *Δget4* and *Δmdy2/get5* tightly clustered with *Δget3* (fig. S5). Additionally, loss of *GET3* fully masked the effect of *Δget4* and *Δmdy2/get5* (Fig. 5A). Moreover, these deletions partially suppressed the UPR induction of *Δget1* and *Δget2*, a phenomenon previously seen with other phenotypes for *Δget3* (32).

Several observations support a role for *GET4* and *MDY2/GET5* in TA protein biogenesis. First, cytosolic extracts from strains lacking Mdy2p/Get5p had a defect in insertion of the model TA substrate

Sec22p into ER microsomes (Fig. 5B). Second, several of the *in vivo* phenotypes characteristic of loss of GET members are also observed in *Δget4* and *Δmdy2/get5* strains. These include a highly significant ($P < 10^{-30}$, Mann-Whitney *U*) relocalization of TA protein GFP-Sed5p from punctate Golgi structures to a more diffuse pattern (Fig. 5C and fig. S7) (12) as well as mislocalization of the peroxisomal TA protein, GFP-Pex15p, to mitochondria (fig. S8) (32). Consistent with a defect in Sed5 biogenesis, loss of Get4p or Mdy2p/Get5p led to secretion of HDEL proteins, a phenotype that is seen in other GET deletion strains (fig. S9). Third, immunoprecipitation revealed that Get4p and Mdy2p/Get5p bind Get3p in the cytosol (Fig. 5D). Mdy2p/Get5p also colocalized with Get3p and TA proteins to punctate protein aggregates that form in *Δget1* strains (32) (figs. S10 and S11). Localization of Get3 to these puncta is dependent on Get4p and Mdy2p/Get5p but not vice versa (figs. S10 to S12), suggesting that Get4p and Mdy2p/Get5p help deliver TA proteins to Get3p in the cytosol for trafficking to the ER membrane. Interestingly, Get4p and Mdy2p/Get5p have been suggested to be peripherally associated with ribosomes (34), where they could potentially capture nascent TA proteins. Thus, whereas Get4p and Mdy2p/Get5p are localized outside of the secretory pathway and initially may have appeared to be false positives, our double mutant analysis revealed how they affect ER protein folding.

Perspective. Our work reveals the range of processes that make the ER a robust folding compartment and yields both a list of components and a blueprint for their functional interdependence. These factors include a wide range of activities such as chaperones, glycosylation enzymes, and ERAD components as well as trafficking pathways, transcriptional regulatory networks, modulators of lipid and ion composition, and vacuolar function. The diversity of activities found supports and extends the recent view in which ER protein folding homeostasis (proteostasis) emerges from the dynamic interplay between folding, degradation, and export processes (35, 36). From a practical perspective, our studies provide a rational starting point for efforts to modulate the ER folding capacity to intervene in disease (36).

More broadly, dissecting complex cellular processes represents a major challenge in cell biology. Deletion libraries and RNA interference (RNAi) approaches now make it possible to identify important factors rapidly (37). But this in turn creates a bottleneck in their functional characterization, which classically requires specialized gene-by-gene follow-up studies. Our approach in effect allows hundreds of different secondary screens to be carried out in parallel to explore systematically the functional interdependencies of hits, thus providing a foundation for focused mechanistic investigations. Given the large number of potential ways of creating proximal reporters for different aspects of biology, our strategy for generalizing systematic quantitative genetic analysis should be broadly applicable to other processes and organisms, including mammals, through the use of double RNAi treatments.

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Supporting Online Material

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References

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