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The Unfolded Protein Response Regulates Multiple Aspects of Secretory and Membrane Protein Biogenesis and Endoplasmic Reticulum Quality Control

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Abstract. The unfolded protein response (UPR) is an intracellular signaling pathway that relays signals from the lumen of the ER to activate target genes in the nucleus. We devised a genetic screen in the yeast Saccharomyces cerevisiae to isolate mutants that are dependent on activation of the pathway for viability. Using this strategy, we isolated mutants affecting various aspects of ER function, including protein translocation, folding, glycosylation, glycosylphosphatidylinositol modification, and ER-associated protein degradation (ERAD). Extending results gleaned from the genetic studies, we demonstrate that the UPR regulates trafficking of proteins at the translocon to balance the needs of biosynthesis and ERAD. The approach also revealed connections of the UPR to other regulatory pathways. In particular, we identified SON1/RPN4, a recently described transcriptional regulator for genes encoding subunits of the proteasome. Our genetic strategy, therefore, offers a powerful means to provide insight into the physiology of the UPR and to identify novel genes with roles in many aspects of secretory and membrane protein biogenesis.

Key words: protein translocation • protein maturation • gene regulation • glycosylation • protein degradation

Introduction

The segregation of specific functions into discrete compartments, or organelles, is a hallmark of all eukaryotic cells. A such, maintenance of organelles and their activities are under precise regulatory control (Nunnari and Walter, 1996). Until recently, little was known regarding the mechanisms used to monitor and respond to a cell’s needs for specific organelle functions. Early clues came from observations of a regulatory pathway in mammalian cells whereby two genes, GRP78 (BiP) and GRP94, are induced by N-linked glycosylation inhibitors or glucose deprivation (Pouyssegur et al., 1977). Since then, perturbations affecting functions of the ER, it was suggested that gene activation was a consequence of ER stress. This notion was demonstrated directly through the expression of protein folding defective mutants of the influenza HA glycoprotein. In cells expressing such mutant proteins, GRP78 (BiP) transcription is induced specifically (Kozutsumi et al., 1988). These early experiments established the existence of a signal transduction pathway between the ER and nucleus, termed the unfolded protein response (UPR) (Mori et al., 1992).

The UPR is a ubiquitous mechanism observed in all eukaryotic organisms from humans to yeast (reviewed in Chapman et al., 1998; Kaufman, 1999). An important step in uncovering the mechanisms underlying the UPR came from promoter studies of the known target gene, KAR2. Deletion analysis revealed specific promoter elements termed UPREs (unfolded protein response elements) that are required for regulated gene activation (Mori et al., 1992; Kohno et al., 1993). The UPRE defined from yeast KAR2 when combined with the CYC1 TATA box is sufficient to drive the UPR-dependent expression of a heterologous reporter gene. This result was key to a genetic strategy for isolating mutants defective for signaling through the UPR (Cox et al., 1993; Mori et al., 1993). The first gene identified encodes an ER transmembrane protein with a cytosol-facing serine/threonine kinase, Ire1p (also termed

Abbreviations used in this paper: CPY, carboxypeptidase Y; ERAD, ER-associated protein degradation; GPI, glycosylphosphatidylinositol; PER, protein processing in the ER gene; UPR, unfolded protein response; UPRE, unfolded protein response element.
E\text{rn1p}), which acts as a sensor of the E\text{R} lumen. Through
an unknown mechanism, stimuli such as accumulation of
misfolded proteins cause Ire1p to oligomerize and trans-
autophosphorylate as a prerequisite step for activation
(Shamu and Walter, 1996). Hereafter, the mechanism of
signal transduction diverges radically from paradigms de-
derived from studies of plasma membrane receptors. U
pon activation, a cytosolic nuclease domain of Ire1p excises an
intronic from HAC1 \text{mRNA} (Cox and Walter, 1996; Kawa-
harata et al., 1997), which encodes a UPR-E-specific tran-
scription activator (Cox and Walter, 1996; Morii et al.,
1996). tRNA ligase then joins the two exons, thereby
completing the splicing of HAC1 \text{mRNA} by a noncon-
ventional, nonspliceosomal mechanism (Sidrauski et al.,
1996). U snspliced HAC1 \text{mRNA} is stable in cells and ini-
tiates translation, but the presence of the intron stalls ribo-
somes so that no Hac1p is produced (Chapman and
Walter, 1997). Removal of the intron upon UPR activa-
tion relieves the translational block to allow synthesis of
Hac1p and the induction of target genes; the Ire1p-medi-
ated splicing reaction, therefore, is a key regulatory step in
the pathway.

Despite the detailed understanding of the mechanisms
surrounding UPR signaling and gene regulation, the pre-
cise physiological role of the pathway has remained largely
unexplored. The most extensive list of targets was assem-
bled from yeast and includes KAR2, LH51, FKBP2, PD11,
EUG1, and ERO1. The products of these genes localize to
the E\text{R} lumen and catalyze protein folding. KAR2 and
LH51 encode proteins with similarity to the Hsp70 class of
molecular chaperones (Normington et al., 1989; Rose et
al., 1989; Craven et al., 1996). FKBP2 is a prolyl isomerase
homologue (Partedalis and Berlin, 1993). PD11, EUG1,
and ERO1 promote disulfide bond formation (Lamanta
et al., 1991; Tachibana and Stevens, 1992; Craven et al.,
1996; Pollard et al., 1998). Thus, the UPR regulates the
abundance of E\text{R} resident chaperones and other enzymes
required for folding, assembly, and modification of secre-
tory and membrane proteins.

The initial identification of Ire1 as a component of
UPR signaling provided additional clues. Ire1 was first
reported as a gene required for inositol prototrophy (Ni-
kawa and Y amashita, 1992). It was later found that regula-
tion of the inositol biosynthetic pathway requires a func-
tional UPR (Cox et al., 1997). The observation showed that
the inositol pathway interacts intimately with the UPR. Since inositol biosynthesis and other aspects of lipid
biosynthesis are coregulated, these observations suggest
that the UPR is involved in the regulation of membrane
biosynthesis. Such a connection may serve to expand the
E\text{R}, when more E\text{R} resident proteins need to be accom-
modated as the result of UPR induction.

Proteins that enter the E\text{R} and cannot be folded cor-
crectly, even after boosting E\text{R} folding capacity through
UPR induction, are degraded. The degradation pathway,
termed E\text{R}-associated protein degradation (ERAD; re-
viewed in Sommer and Wolf, 1997; Brodsky and Mccrak-
en, 1999), translocates misfolded proteins back into the
cytosol, where they are degraded by the proteasome.
Retrotranslocation (also called dislocation) is thought to
utilize the same core protein complex (Sec61p and associ-
ated subunits) that forms the protein conducting channel
in the translocon through which proteins are delivered to
the E\text{R} lumen. Conceptually, the UPR, in its previously
known scope, and ERAD provide different means of dealing with
protein misfolding in the E\text{R}: the UPR by inducing enzymes
thought to play a corrective role and ERAD to dispose of
proteins that cannot be rescued. Here, and in a concomitant
study (Travers et al., 2000), we show that the two pathways
indeed are intimately linked and that the scope of the UPR
encompasses many more aspects of protein maturation and
ER quality control than previously appreciated.

Materials and Methods

Strains and Antibodies

Yeast strains used in this study are described in Table I. A nticarboxypeptidase Y (anti-CPY) antisera generously provided by Dr. Reid Gilmere
(University of Massachusetts, Worcester, MA). A nti-Gas1p antisera
was a kind gift of Dr. Howard Riezman (University of Basel, Switzerland).
A nti-HA mAb (12CA5) was generated by Berkeley A ntibody Company.

Plasmids Used in This Study

pCS15 (SEC61, LEU2) was provided by Dr. Randy Schekman (University of California, Berkeley, CA) and pMR713 (KAR2, LEU2) was provided by Dr. M ark Rose (Princeton University, Princeton, NJ).

Construction of pDN336 and pDN388. The plasmid pDN336 used as the reporter in strain DNY421 was constructed by inserting the full-length
Ire1p gene, released from pCS110 (Cox et al., 1993) as an X hoI/BamHI
fragment, and the full-length Ade3 gene, released as a BamHI/Bhel fragment,
into X hoI/XbaI sites of the yeast shuttle vector, pRS316 (URA3,
CEN6, ARSH4; Sikorski and H ieter, 1989). Both the XbaI and the Nhel sites
were destroyed in the construction. pDN388 is similar, except that the
insert is in the pRS315 vector (LEU2, CEN6, ARSH4; Sikorski and
H ieter, 1989).

pDN390. The plasmid, pC835, containing the HAC1 gene in the shut-
tle vector, pRS313 (Cox and Walter, 1996), was digested with A wni
and NgotI to release the gene. The fragment was ligated into pRS313
generated with the same sites.

CPY\text{HA} Expression Vectors. The prc1-1 allele expresses the variant
CPY\text{HA} as a glycine to arginine change at position 255 (Finger et al., 1993).
We constructed an HA epitope-tagged version of CPY\text{HA} by site-directed
mutagenesis using a PCR-based approach. The PRCl gene was first am-
plified as two fragments. The N-proximal fragment amplified with the
primers 5'-CCATCGAATTCGCTATGATG-3' and the phosphorylated
primer 5'-GAAATCTTGGCCGGTTGACG-3' using Vent polymer-
ase (New England Biolabs). This fragment includes the PRCl promoter
and coding sequences to amino acid position 251. C-proximal sequences
were amplified using the phosphorylated primer 5'-CAATCTGCTA-
GAAATCTTGGCCGGTTGACG-3' incorporating a glycine to arginine change
at amino acid 251 and the primer 5'-CCCTCTAGACTAACCCAAAG-
AAGGATTAACCTGGTAACTG-3', fusing the HA epitope tag, a termination codon, and an XbaI
site. The C-proximal fragment was amplified using Taq polymerase since
the primer combination precluded amplification by Vent polymerase un-
der all conditions attempted. Blunt ends were generated for this fragment
using T4 DNA polymerase. The N-proximal fragment was digested with
EcoR1, the C-proximal with XbaI, and both fragments ligated into the
vector pDN201 (Ng et al., 1996) digested with same enzymes. The mutant
version of the plasmid, pDN436, was constructed by releasing the CPY\text{HA} dad
gene from pDN431 as a SalI/EcoR1 (blunt) fragment and ligation into the
SalI/Smal sites of pRS313 (Sikorski and H ieter, 1989).

Genetic Screen

The reporter strain for the screen was constructed by first crossing JC147
and EY0060. Diploids derived were sporulated, tetrads dissected, and
haploids screened for the following genotype:

\text{M A T a } \text{i r e 1:: T R P 1 , a d e 2 ,}
\text{a d e 3 , U F P R - L a c Z:: H I S 3 . O n e such strain was isolated and transformed}
with pDN336 to create DNY421.

For mutagenesis, 50 A1000 OD units of DNY421 cells were washed once

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in sterile water and resuspended in 25 ml 0.9% KCl. With constant stirring, the cells were grown for 15 cm above. After each pulse, an aliquot was removed, a portion serially diluted, and spread onto YPD plates to determine the kill rate. The remainder of each aliquot was pelleted and resuspended in 10 ml YPD media and allowed to recover for 18 h on a roller drum. All procedures to this point were performed in a darkroom and incubations at 30°C. Cells receiving a UV dose of 100 J/m² resulted in a kill rate of 62% and were used for the screen. Mutagenized cells were spread onto YPD plates lacking additional adenine. Nonsectoring colonies were picked and rescreened using additional adenine. Nonsectoring colonies were picked and rescreened by restreaking for single colonies (see Results for specific numbers). Iso-

| Strain | Genotype | Source |
|--------|----------|--------|
| W303a  | MATa, leu2-3,112, his3-11, trp1-1, ura3-1, can1-100, ade2-1 | Walter lab |
| EY0860 | MATa, ade3, W303 background | Cox et al., 1993 |
| JC17   | ire1::TRP1, ura3-1, can1-100, ade2-1, leu2-3,112::LEU2-UPRE LacZ, his3-11::HIS3-UPRE LacZ | Cox et al., 1993 |
| JC408  | MATa, lac2::URA3, trp1-1, his3-11, ade2-1, leu2-3,112::LEU2-UPRE LacZ | Walter lab |
| JC409  | MATa, JC408 background | Walter lab |
| DNY70  | sec62-101, ura3Δ99, leu2Δ1, trp1Δ99, ade2-103 | Ng et al., 1996 |
| DNY420 | ire1::TRP1, ura3-1, can1-100, ade2-1, ade3, leu2-3,112::LEU2-UPRE LacZ, his3-11::HIS3-UPRE LacZ, (pDN336) | This study |
| DNY421 | ire1::TRP1, ura3-1, can1-100, ade2-1, ade3, leu2-3,112::HIS3-UPRE LacZ, (pDN336) | This study |
| DNY486 | per1-1, DNY421 background | This study |
| DNY499 | per1-2, DNY421 background | This study |
| DNY498 | per2-1, DNY421 background | This study |
| DNY491 | per3-1, DNY421 background | This study |
| DNY493 | per4-1, DNY421 background | This study |
| DNY478 | per4-2, DNY421 background | This study |
| DNY495 | per5-1, DNY421 background | This study |
| DNY497 | per6-1, DNY421 background | This study |
| DNY503 | per6-2, DNY421 background | This study |
| DNY501 | per7-1, DNY421 background | This study |
| DNY505 | per8-1, DNY421 background | This study |
| DNY507 | per9-1, DNY421 background | This study |
| DNY509 | per10-1, DNY421 background | This study |
| DNY484 | per11-2, DNY421 background | This study |
| DNY488 | per11-1, DNY421 background | This study |
| DNY470 | per12-1, DNY421 background | This study |
| DNY472 | per13-1, DNY421 background | This study |
| DNY475 | per14-1, DNY421 background | This study |
| DNY479 | per15-1, DNY421 background | This study |
| DNY481 | per16-1, DNY421 background | This study |
| DNY523 | per1-1, W303 background | This study |
| DNY540 | per15-1, W303 background | This study |
| DNY563 | pDN431, W303 background | This study |
| DNY572 | macl::TRP1, pDN431, W303 background | This study |
| ESY157 | ire1::TRP1, pDN431, W303 background | This study |
| ESY158 | ESY157 with pMR713 (KAR2) | This study |
| ESY159 | ESY157 with pCS15 (SEC61) | This study |

Cloning and Identification of PER Genes

Two general approaches were employed to clone PER (protein processing in the ER) genes. The first takes advantage of the counterselectable marker, URA3, contained in the pDN336 (Boeke et al., 1984). Since the growth of per mutants is dependent on the Ire1p, cells losing pDN336 fail to grow on media containing 5-fluoroorotic acid (5-FOA). Complementation of mutant alleles alleviates the requirement for Ire1p and thus allows for selection of complementing plasmids from genomic libraries. The PER5 and PER8 genes were cloned using this first approach.

Mutant cells were transformed with a yeast genomic library based on the multicopy YEp13 vector (Lagodzky et al., 1987). A few transformations, the cells were grown overnight in SC-Leu media allowing transformants receiving complementing plasmids to lose pDN336. Transformants were later plated and incubated on SC-Leu media containing 5-FOA (1 mg/ml) at 30°C. Plasmids were recovered from 5-FOA-resistant isolates by zirconium bead disruption and purification using the Wizard MiniPrep kit (Promega). Plasmid amplification was performed after transformation into bacterial DH5α cells. Retransformation of the recovered plasmids into the respective mutant strains was typically carried out to assess complementation of the sectoring phenotypes. A through this approach worked well for PER5 and PER8, but was less successful for other strains attempted. The high incidence of false positives due to plasmids carrying truncated Ire1p in the library added an additional layer to the procedure that was overly time consuming.

The second approach scored for complementation of the sectoring phenotype using a low copy genomic library (used for cloning of PER2, PER4, PER13, and PER16). The library, based on YCP50 (Rose et al., 1987), required that the reporter pDN336 be swapped for pDN38 to be compatible. Mutant strains transformed by the library were spread onto SC-Ura/adenine-limiting (6 µg/ml) plates and incubated at 30°C. Since it was less successful for other strains tested. The high incidence of false positives due to plasmids carrying truncated Ire1p contained in the library added an additional layer to the procedure that was overly time consuming.

The second approach scored for complementation of the sectoring phenotype using a low copy genomic library (used for cloning of PER2, PER4, PER13, and PER16). The library, based on YCP50 (Rose et al., 1987), required that the reporter pDN336 be swapped for pDN38 to be compatible. Mutant strains transformed by the library were spread onto SC-Ura/adenine-limiting (6 µg/ml) plates at low density (400 cfu/plate) to develop the colony color phenotype. Typically, between 10,000 and 25,000 transformants were screened for restoration of sectoring. Positives were cured of the reporter and complementing plasmids were recovered as described above. Recovered plasmids were transformed back into the respective mutant strains to confirm complementation.
For all clones, DNA sequence analysis was performed to determine the identity of inserts (Nucleic Acid Facility, Pennsylvania State University, University Park, PA). The sequences of vector/insert junctions were obtained using the primers N168 (5'-CGCTACTTGAGCCGCACTATCGAC-3') and N169 (5'-ATGGTCTATCGCCGATAT-3'). Junction sequences were submitted to the Saccharomyces Genome Database (http://genome-www.stanford.edu/Saccharomyces) to obtain complete insert sequences and identities of open reading frames. The insert sequences were used to facilitate deletion mapping using standard recombinant DNA methods to determine specific complementing genes. Subclones of genes derived from the high copy library (YEp13) were inserted into centromeric plasmids to assess complementation.

Cell Labeling and Immunoprecipitation

Typically, 3 A od units of log phase cells were pelleted and resuspended in 0.9 ml of SC media lacking methionine and cysteine. A sterile centrifugation, the TCA precipitate pellet was resuspended in 120 ml of IPS II (100 mM Tris base, 3% SDS, 1 mM PM SF) and heated to 100°C for 5 min. Insoluble debris was pelleted and 40 ml of the detergent lysate was added to 560 ml IPS II (1% Triton X-100, 50 mM Tris, pH 7.5, 1 mM PM SF, 1 ml yeast protease inhibitor cocktail; Sigma-Aldrich) and the appropriate antiserum. A sterile 2-h incubation at 4°C, the sample was centrifuged for 10 min at 16,000 g and the supernatant transferred to a fresh tube containing protein A-Sepharose beads. The tube was resuspended for 30 min and washed three to five times with IPS I (0.2% SDS, 1% Triton X-100, 50 mM Tris, pH 7.5) and once with PBS. Immunoprecipitated proteins were eluted with gel sample buffer, separated by gel electrophoresis, and visualized by autoradiography. In pulse-chase assays to determine protein stability, immunoprecipitations were normalized by measuring TCA-precipitable counts using an LS801 scintillation counter (Beckman Coulter) and adjusting lysates for equal counts.

Northern Blot Analysis

Preparation of RNA, gel electrophoresis, blot transfer, and probing were performed as described in Cox and Walter (1996). RNA from UPR-induced cells were treated for 60 min with 2.5 μg/ml tunicamycin (Sigma-Aldrich) before RNA isolation. Two templates for probes were prepared by PCR amplification of genomic DNA. Probes were prepared by random primer extension labeling using [TPP]-aCTP. The ACT1 template is a 600-bp fragment corresponding to the 3’-end of coding sequences. The KAR2 template is a 600-bp fragment corresponding to the 5’-end of coding sequences. Two RFT1 templates were prepared and used independently. One includes 600 bp of the 3’-coding sequences and the other includes 700 bp of the 3’-coding sequences.

Results

Genetic Screen

We devised two complementary approaches to identify in an unbiased way cell functions that are regulated by or are dependent on the UPR. First, we used whole genome microarray analysis to directly identify UPR target genes (Travers et al., 2000). This approach allowed us to define the transcriptional scope of the UPR. Second, we devised the genetic screen that is described here in which we isolated mutants that are dependent on UPR activation for viability. This second approach aims to highlight physiologically relevant functions that connect with the UPR. The feasibility of this approach was suggested by the observations that genes mediating the UPR are nonessential (Cox et al., 1993; Mori et al., 1993), yet display synthetic lethality (that is, cell inviability caused by the combination of two nonlethal mutations) with two known target genes (KAR2 and LHS1; Craven et al., 1996; Sidrauski et al., 1996). In both kar2 and lhs1 mutants, the UPR is constitutively activated, which must allow the cell to compensate for the loss of the function of these genes. In principle, genetic and microarray analyses should be complementary, as some functions that are dependent on or affected by UPR activation may not be transcriptional targets of the pathway.

We performed the screen using a yeast colony color sorting assay (Koshland et al., 1985). Yeast cells harboring an ade2 mutation give rise to red colonies when grown on media limiting for adenine. If a strain contains both mutants ade2 and ade3 alleles, the red pigment is not synthesized, and colonies are white. We constructed an ade2 ade3 mutant strain that is deleted for IRE1 (DNY421, Table I). This strain grows well on rich media. In addition, we constructed a centromeric reporter plasmid, pDN336, containing IRE1 and ADE3, which was transformed into DNY421 cells. Since ADE3 and IRE1 are not essential, the plasmid is lost at a frequency of ~10^-6 per cell division (Guthrie and Fink, 1991). For this reason, DNY421 cells give rise to red and white sectored colonies, reflecting a mixed population of cells with and without the plasmid. Because IRE1 is required for activation of the UPR pathway, mutants that require UPR activation for growth cannot lose the IRE1-bearing plasmid and hence, are expected to give rise to red nonsectored colonies.

DNY421 cells were mutagenized with ultraviolet light to 62% lethality. We screened ~50,000 colonies and isolated 104 nonsectoring and normally growing colonies. Of these, 40 were backcrossed to the parental strain, yielding 32 re-cessive, 2 dominant, and 6 sterile isolates. The dominant and sterile mutants were discarded. Heterozygous diploids generated from recessive mutants were sporulated and subjected to tetrad analysis to determine the segregation patterns. Of the diploids, 25 were sporulation-competent. Of those, 15 segregated 2:2 with a synthetic lethal phenotype and 5 with a synthetic negative growth phenotype (spores containing both mutations were viable, but displayed severely reduced growth rates), indicative of a single gene being responsible for the mutant phenotype. Of the remaining mutants, five exhibited other patterns and were discarded.

When crossed to each other, many of the mutants exhibited partial sectoring phenotypes that precluded clear determination of complementation. This was not a general effect of ploidy since all 20 mutants, when crossed to the parental strain, formed sectored colonies unambiguously. Further analysis demonstrated that most mutant genes are unlinked. These observations suggest some combinations of mutations display the genetic interaction termed unlinked noncomplementation. Although the significance for this group remains to be determined, unlinked noncomplementation has been used to help identify genes of the same function or pathway (Stearns and Botstein, 1988). To group the 20 mutants, we crossed individual mutant strains and systematically subjected the resulting diploids to tetrad analysis. Most crosses failed to display linkage between the mutant alleles, indicating that the mutations
were in different genes. This analysis defined 16 linkage groups indicating at least as many genes (Table I). We refer to the genes defined by the mutants as PER (protein processing in the ER) because most mutant alleles can be shown to affect various aspects of ER protein biogenesis and quality control (see below). The high fraction of per mutants represented by a single allele indicates that the screen is far from saturated.

To establish PER mutants as tools to study the functional role of the UPR, we next assessed their physiological relevance. To this end, we required two criteria to be met: activation of the UPR pathway should alleviate the synthetic growth defect caused by the mutation, and the mutants should exhibit a constitutive activation of the UPR. To address the first criterion, we activated the UPR constitutively by expression of the Hac1p transcriptional activator (Cox and Walter, 1996). We constructed a plasmid, pDN390, carrying HAC1 (HAC1 deleted of its intron) and transformed it into each per mutant (ΔIRE1 cells covered by pDN336 bearing wild-type IRE1). Cells were then scored for improved growth after loss of pDN336, obtained by screening for white, nonsectoring colonies. Using these criteria, all but two mutants (per1-1 and per15-1) could be relieved of the requirement for IRE1 function by directly activating the pathway (Table I). A fter crosses of the two remaining mutants (DNY523 and DNY540) with Δhac1 cells were sporulated, however, tetrad analysis revealed clear synthetic negative phenotypes for both mutants (data not shown). Thus, taken together, the results indicate that the synthetic defect of all 16 per mutants is due to the loss of the UPR pathway, rather than to the loss of a yet uncharacterized function of IRE1 unrelated to UPR signaling.

To address the second criterion for physiological relevance, we determined the extent of constitutive UPR activation in mutant cells. The parental strain (DNY421) used in this study contains a genomic copy of a lacZ reporter gene fused to a minimal UPR promoter (Cox et al., 1993). Thus, ß-galactosidase activity can be used to monitor activation of the pathway. Mutant cells were grown to early log phase, and cell extracts were prepared to measure ß-galactosidase activity with the substrate 2-nitrophenyl-ß-D-galactopyranoside (Cox et al., 1993). As shown in Fig. 1, with the exception of per16-1, all per mutants express ß-galactosidase activity above wild-type levels, indicating induction of the UPR. Interestingly, the degree of UPR induction among the mutants varies widely, indicating that the UPR can be gradually modulated depending on the physiological needs of the cell.

### Functions of PER Genes

Having satisfied the criteria for a physiological relationship, we next assessed functional defects of per mutants. To this end, we first monitored the processing and transport of the membrane protein Gas1p (Nuoffer et al., 1991). Because Gas1p undergoes a variety of posttranslational modifications during its biogenesis, defects at any stage can be detected by changes in gel mobility. Gas1p is initially synthesized in the cytosol as a 60-kD precursor that is detectable in protein translocation mutants (Ng et al., 1996). Upon entry into the ER, it is modified by N- and O-linked glycosylation, as well as by addition of a glycosylphosphatidylinositol (GPI) membrane anchor shifting its apparent molecular weight to 110 kD (Nuoffer et al., 1991). Gas1p folding occurs in the ER and requires the formation of intramolecular disulfide bonds (Frand and Kaiser, 1998). Only after correct folding can Gas1p continue to the Golgi apparatus (t½ < 10 min), where carbohydrate modification further changes its gel mobility to
Thus, defects in a variety of ER functions can be detected by monitoring Gas1p processing.

Wild-type (W303) and per mutant cells were metabolically pulse-labeled with [35S]amino acids for five minutes and chased with an excess of unlabeled methionine and cysteine for 30 min to allow for full processing of Gas1p (Fig. 2 A, lanes 2–22). As shown in Fig. 2, no preGas1p was observed in any lane, indicating that none of the per mutants displayed severe defects in protein translocation. However, about half of the mutants are defective in processing to the mature 125 kD form. These results indicate defects in ER protein processing functions that may include glycosylation, folding, GPI-anchor addition, or transport.

To distinguish the molecular nature of the observed defects further, we monitored the processing of another protein, the vacuolar protease CPY (Simons et al., 1995). The ER form of CPY is a core glycosylated proform (P1; Fig. 2 B, lane 1). The P1 form of CPY is transported to the Golgi apparatus, where it is modified by outer chain glycosylation to the P2 form. ProCPY is ultimately proteolytically processed to the mature form after transport to the vacuole (Fig. 2 B, lane 2). Underglycosylated CPY remains competent for folding and transport to the Golgi apparatus and vacuole. In the vacuole, it is processed to characteristic forms designated -1, -2, -3, -4 (te Hesen et al., 1992). Thus, based on specific gel mobility patterns, mutants defective for N-linked glycosylation can be easily distinguished.

As shown in Fig. 2 B, mutants per3-1, per5-1, per6-1, per6-2, per12-1, and per14-1 exhibited CPY underglycosylation indicative of defective N-linked glycosylation. Correspondingly, these mutants synthesized forms of Gas1p with altered mobility expected for glycosylation mutants (Fig. 2 A). The size of the underglycosylated forms indicates that most of CPY is proteolytically processed to the mature form in each of the mutants, indicating that these mutants are not generally defective in protein transport from the ER to the Golgi apparatus.

The PER5/RFT1 Gene Is a Novel UPR Target Required for N-linked Glycosylation

We chose to analyze per5-1 in greater detail because it exhibited the most severe glycosylation defect. To this end, we analyzed CPY by pulse-chase analysis, followed by en...
doglycosidase H digestion in wild-type and per5-1 cells. As shown in Fig. 3 A, proCPY recovered after the pulse migrated as multiple species with altered mobility in per5-1 cells as compared with the P1 species from wild-type cells (Fig. 3 A, compare lanes 1 and 4). Removal of the sugar chains revealed that the heterogeneity was exclusively due to differences in glycosylation, as the deglycosylated forms comigrated (Fig. 3 A, lanes 7–12). The kinetics of processing to the mature form(s) was similar in both strains, indicating that folding and transport functions are intact in per5-1 cells. Similar results were obtained for Gas1p (Fig. 3 B), indicating that this defect is not substrate-specific.

To characterize the role of PER5 further, we cloned the gene by complementation (see Materials and Methods). A complementing plasmid, pDN386, was isolated from transformed per5-1 cells and portions of the insert further subcloned to identify the complementing gene. Plasmid pDN387, containing RFT1 as the only open reading frame, fully complemented per5-1 as it restored the sectoring phenotype and glycosylation function (Fig. 3 B). RFT1 is an essential gene that encodes a predicted multispanning transmembrane protein of unknown function (Koerte et al., 1995). Thus, our approach identified PER5/RFT1 as a novel gene required for N-linked glycosylation.

Inspection of the upstream sequences of PER5/RFT1 revealed a potential UPRE, suggesting that transcription of PER5 is regulated by the UPR (Fig. 4 A). To address this possibility directly, Northern analysis was performed using RNA extracted from control cells and from cells treated with the glycosylation inhibitor tunicamycin to induce UPR-regulated genes. As shown in Fig. 4 B, PER5/RFT1 is elevated in tunicamycin-treated cells (lanes 1–4). Quantitative analysis showed a moderate 2.5-fold induction. This is in contrast to KAR2, which was induced over 8-fold (Fig. 4 B). Transcriptional regulation of PER5/RFT1, as with KAR2, was dependent on the UPR, as no induction was detected in Δhac1 cells (Fig. 4 B, lanes 5 and 6).

Taken together, the analyses in Figs. 2–4 show that mutations in at least 5 of 16 PER genes compromise important functions in protein glycosylation and that genes of heretofore unknown function have been identified in this way. Moreover, these results reinforce the notion that the UPR functions to regulate a variety of aspects of protein biogenesis.

A Role for the UPR in ERAD

As many of the per mutants do not exhibit defects in Gas1p or CPY maturation, we wondered whether other aspects of ER function related to protein maturation are affected in some of the per mutants. To test whether
genes encoding ERAD components were identified in this screen, we constructed a recombinant form of the ERAD substrate, CPY* (Finger et al., 1993), modified by addition of an HA epitope tag to simplify analysis. The new version, designated CPY*HA, behaved as a proper ERAD substrate: it was degraded rapidly in wild-type cells and was stabilized in a strain lacking the ERAD gene, CUE1 (Fig. 5; Biederer et al., 1997). We transformed each of the per mutants with pDN436, carrying the gene encoding CPY*HA. Pulse-chase analysis was used to assess substrate stability in each strain. Turnover of CPY*HA was delayed in per7-1, per8-1, per9-1, per13-1, and per16-1 (Fig. 5). Two mutants, per9-1 and per13-1, also displayed defects in Gas1p processing, indicating that their defects are pleiotropic. Their ERAD phenotypes may therefore be indirect. In all other per strains, including those displaying severely impaired protein processing, the rate of CPY*HA degradation was not altered (data not shown).

We next cloned some of the affected genes of per mutants displaying ERAD defects, starting with PER8 and PER16, which displayed defects similar in extent to the Δcue1 strain. Plasmid pDN426 bearing SON1, completely complemented the per8-1 mutant. Moreover, CPY*HA was stabilized in αson1 cells (data not shown), indicating that the defect per8-1 cells reflects a loss-of-function. Son1p was recently shown to be a transcription factor that regulates proteasome biogenesis (Mannhaupt et al., 1999), and Δson1 mutants are defective in cytosolic protein degrada-

Figure 5. A subset of per mutants is defective for ERAD. All twenty per mutants were transformed with pDN436 (CPY*HA) and pulse-chase analysis performed to measure the degradation of the ERAD substrate CPY*HA. Immunoprecipitates of CPY*HA were normalized using TCA precipitable counts. The proteins were separated by PAGE followed by autoradiography. Quantification was performed by PhosphorImager analysis and reported as percent remaining to the right of each respective autoradiogram. In addition to the wild-type (DNY 563) and ERAD mutant (DNY 572, cue1::TRP1) controls, only per mutants with significant ERAD defects are shown.

Figure 6. The UPR is required for efficient ER protein translocation and ERAD. A, Wild-type and ire1::TRP1 cells (labeled Δire1) expressing CPY*HA were pulse-labeled for 10 min with [35S]methionine/cysteine and chased for 0, 30, 60, and 90 min. CPY*HA was immunoprecipitated using anti-HA mAb and analyzed by SDS-PAGE. The proteins were visualized by direct autoradiography and quantification of the gel was performed using a phosphorimager. B, Wild-type, sec62-101, and Δire1 cells expressing CPY*HA were pulse-labeled for 10 min and CPY*HA immunoprecipitated and separated on a 10% polyacrylamide gel (lanes 1, 2, and 3). Lanes 4 and 5, Gas1p immunoprecipitated from the lysates used for lanes 1 and 3. C, Wild-type, sec62-101, and Δire1 cells expressing CPY*HA were pulse-labeled for 10 min and immunoprecipitated for endogenous CPY. The position of preproCPY·HA, the ER P1, and Golgi P2 forms are indicated.

We cloned PER16 by screening for restoration of the sectoring phenotype after a plasmid shuffle of the URA3-marked reporter for LEU2 (pDN 388) and transformation of the library. Subcloning of the inserts of plasmids that restored the sectoring phenotype of per16-1 cells identified UBC7, which was sufficient to complement the per16-1 mutation. UBC7 encodes a ubiquitin-conjugating enzyme that was previously shown to participate in ERAD (Biederer et al., 1996; Hiller et al., 1996).

The isolation of ERAD mutants in this screen suggested a tight physiological link between the UPR and ERAD. We explored this notion further by monitoring the fate of CPY*HA in a UPR-deficient strain. As shown in Fig. 6 A,
CPY*HA was stabilized in Δire1 cells (Fig. 6 A, lanes 5–8), whereas wild-type control cells degraded the substrate rapidly. Surprisingly, we also observed the appearance of a faster migrating band in Δire1 cells that was not detected in control experiments performed with wild-type cells (Fig. 6 A, labeled preproCPY*HA). Its mobility and the timing of its appearance during the pulse suggested that the band represents the cytoplasmic, nontranslocated preproCPY*HA. To confirm this identity, we expressed CPY*HA in sec62-101 cells, in which translocation of CPY is severely impaired and thus accumulated cytosolic precursor proteins. The data in Fig. 6 B show that the preproCPY bands comigrate (compare lanes 2 and 3).

These results raised the question whether impaired translocation resulted from the synthesis of misfolded protein or from an increased flux of proteins through the ER translocon due to ectopic expression of CPY*HA. To distinguish between these possibilities, we constructed wild-type and Δire1 strains to express epitope-tagged wild-type CPY (CPYHA) using the same vector as for CPY*HA. Expression of CPYHA caused no translocation defects in Δire1 cells (Fig. 6 C), indicating that the defect results specifically from the expression of a misfolded protein that is a substrate of the ERAD pathway.

In Δire1 cells expressing CPY*HA, we also observed a translocation defect for the endogenous protein Gas1p (Fig. 6 B, lanes 4 and 5), suggesting that the misfolded protein causes a general translocation defect in these cells. By contrast, Δire1 cells alone or Δire1 cells expressing CPYHA exhibit no defects in ER import of Gas1p (data not shown). A protein import into and export from the ER are thought to share the same translocation pore, it is possible that both processes directly compete for some limiting component(s) regulated by the UPR.

Quantitative analysis of the data in Fig. 6 A showed a delay of ER protein degradation in Δire1 cells. In wild-type cells, 47, 13, and 3% of CPY*HA remained after 30, 60, and 90 minutes of chase, respectively. By contrast, in Δire1 cells, 46, 37, and 22% of CPY*HA remained at the corresponding time points. This delay could result from two, possibly additive, effects: an impairment of import into the ER and a delay in reexport from the ER for degradation. Although the relative contributions of the two processes to the overall delay remains to be determined, ERAD function remains disrupted if the import block is alleviated (see below).

Taken together, the data suggest a role of the UPR in balancing the trafficking of proteins into and out of the ER during periods of stress as import and export substrates compete for a limiting core translocation machinery. The accumulation of CPY*HA would induce the UPR to augment the translocation machinery to clear the ER of the misfolded proteins while maintaining protein import into the ER. In support of this hypothesis, genes encoding components of the translocation machinery with roles for both import and export are targets of the UPR (Travers et al., 2000). In addition, CPY* and CPY*HA expression each cause UPR induction, albeit at a modest level of less than twofold as measured by a UPRE-lacZ reporter (Knop et al., 1996; Spear, E.D., and D.T.W. Ng, unpublished results).

To test this hypothesis, we assessed the contribution of two specific UPR target genes by increasing their expression as it would occur during normal UPR activation. To this end, we tested strains that contain duplicated genes encoding Kar2p or Sec61p, respectively. These genes were selected because of their dual roles in both ER protein import and ERAD (Pilon et al., 1997; Plemper et al., 1997; Zhou and Schekman, 1999). In addition, both genes are inducible under conditions that activate the UPR (Travers et al., 2000). As shown in Fig. 7 A, increased gene dosage of either Kar2p or Sec61p was sufficient to alleviate the translocation defect for preproCPY*HA and preGas1p in Δire1 cells. Δire1 cells with increased expression of Kar2p also alleviated the delay in CPY*HA degradation (Fig. 7, B and C). In contrast, Δire1 cells containing an additional copy of SEC61 remain defective in degrading CPY*HA. The additional copy of SEC61 itself is not inhibitory to ERAD since wild-type cells under the same circumstances show normal if slightly accelerated ERAD function (Spear, E.D., and D.T.W. Ng, unpublished results). These results show that Kar2p becomes limiting for both translocation
and ERAD when cells are challenged with CPY\textsuperscript{*HA}, whereas Sec61p becomes limiting only for translocation, but not ERAD. Moreover, increased expression of either Kar2p or Sec61p can compensate for the translocation defect in \textit{Aire1} cells. Taken together, these data demonstrate an important role for the UPR in regulating the import and export functions across the ER membrane.

**The UPR Controls a Wide Variety of ER Functions**

As described so far, the genetic approach has revealed two new functions, ERAD and glycosylation, that, when limiting, require activation of the unfolded protein response for cell viability. In addition, cloning of \textit{per13} and \textit{per2} revealed that the mutations map to \textit{GPI10} and \textit{MCD4}, respectively, both encoding essential components in GPI anchor addition. A \textit{MCD4}, other genes required for GPI biosynthesis, including \textit{GAA1}, \textit{GPI12}, and \textit{LAS21} are also regulated by the UPR (Travers et al., 2000); \textit{GPI10}, in contrast, was not identified as a UPR transcriptional target gene. Finally, cloning of \textit{per4} revealed that the mutation resides in \textit{LHS1/SSI1}, an hsp70-like ER chaperone (Baxter et al., 1996; Craven et al., 1996), thus adding to the list of classical UPR targets that mediate protein folding directly. Consistent with previous observations, the \textit{per4} mutants display a minor protein translocation defect following a short pulse label (data not shown). Thus, genetic analysis reveals a wide variety of ER functions that are physiologically linked to the UPR.

**Discussion**

Characterization of the \textit{per} mutants revealed that cells require a functional UPR to cope with and compensate for defects in many aspects of protein maturation in the ER, including protein folding, glycosylation, and GPI anchor addition. In addition, we found that ERAD, required to clear the ER of unwanted proteins, is intimately linked to the UPR. A gene expression chip analysis performed in a parallel study to determine the transcriptional scope of the UPR revealed that over 350 genes are transcriptionally upregulated upon induction of the pathway (Travers et al., 2000). Many of the identified genes function in various aspects of the secretory pathway at the level of the ER or beyond. Thus, together the results of the two studies show that the role of the UPR is multifaceted and much more complex than previously appreciated. Importantly, not all of the genes identified as \textit{per} mutants are transcriptional targets of the UPR. Genetic and gene expression array analyses therefore proved complementary rather than redundant approaches, with both avenues allowing the identification of novel genes that function in these processes.

**Regulation of Protein Glycosylation**

Isolation of the \textit{per5}-1 mutant led to the discovery of a new gene involved in protein glycosylation. We found that \textit{PER5} is upregulated by the UPR. \textit{PER5/RFT1} was originally cloned from a yeast mutant requiring the overexpression of human p53 for viability (Koerte et al., 1995); as there is no yeast equivalent to mammalian p53, the reason for this phenotype and the function of \textit{PER5/RFT1} has remained obscure. Our studies demonstrate that \textit{PER5/RFT1} is required for efficient N-linked glycosylation of glycoproteins. \textit{PER5/RFT1} encodes a multispansing transmembrane protein that bears no significant sequence similarity to any other known component of the glycosylation machinery. Our results suggest that \textit{PER5/RFT1} functions in biosynthesis of the dolichol-linked carbohydrate precursor or in the protein conjugation step. Preliminary experiments indicate that \textit{per5}-1 cells show an accumulation of a glycosylated intermediate (Dol-PP-GlcNAc\textsubscript{2}-M\textsubscript{an3}) and a sharp decrease in the mature form (Dol-PP-GlcNAc\textsubscript{2}-M\textsubscript{an3}Glc3; J. Helenius, D.T.W. Ng, P. Walter, and M. Aebi, unpublished results). These results suggest the intriguing possibility that \textit{PER5/RFT1} may encode the long-sought-after flippase that translocates the Dol-PP-GlcNAc\textsubscript{2}-M\textsubscript{an3} intermediate (which is synthesized on the cytosolic face of the membrane) to the lumenal face for further processing (Burd and Aebi, 1999). Consistent with this view, those carbohydrates that are attached to proteins in \textit{per5}-1 cells were found to be fully mature. Further experiments to address this notion are in progress.

Genome expression data has shown that additional components of the glycosylation biosynthetic machinery are regulated by the UPR (Travers et al., 2000). The regulation of glycosylation components helps cement our broadened view of the UPR, as it indicates a role of the pathway in adjusting the cell’s biosynthetic capacity of the ER according to need. Similarly, we isolated per mutants defective in GPI anchorage, and many GPI biosynthetic genes are UPR targets (Travers et al., 2000).

**UPR Regulation of ER Quality Control**

The identification of ERAD genes suggested a tight physiological link between ERAD and the UPR. We have provided direct evidence for such a link using CPY\textsuperscript{*HA}, which is stabilized in UPR-deficient cells. Consistent with previous studies (Kawahara et al., 1997; Zhou and Schekman, 1999), we show that various ERAD-deficient mutants constitutively induce the UPR. As shown in the parallel study, most known ERAD genes are activated by the UPR (Travers et al., 2000). Moreover, in this study we have shown that UPR induction increases ERAD efficiency. Thus the UPR and ERAD are intimately coordinated and interdependent.

The identification of \textit{SON1} (\textit{PER8}) as a gene required for ERAD provided an unexpected twist to the outcome of our genetic screen. \textit{SON1} (also called \textit{RPN4}) was recently shown to be a transcriptional activator that binds a consensus promoter element, termed PACE (Manhart et al., 1999). The PACE motif is found in many genes, but is a common element in the family of genes encoding subunits of the 26S proteasome. In \textit{son1} null mutants, the degradation of some cytosolic proteins are impaired at a step following ubiquitin conjugation (Johnson et al., 1995). Together, these data suggest \textit{SON1} regulates aspects of proteasome biogenesis, which in turn is required for ERAD. Surprisingly however, neither \textit{SON1} nor proteasomal subunit genes are regulated by the UPR (Travers et al., 2000). Rather, proteasomal subunit genes are coordinately upregulated in UPR-deficient cells following treatment with tunicamycin or DTT (Travers et al., 2000). These results suggest that, in addition to the UPR, a UPR-independent
ER → nucleus signal transduction pathway exists in which Son1p may play an important role. Interestingly, the UPR target genes PDI1 and PER5/RFT1 contain consensus PA CE promoter elements (Mannhaupt et al., 1999), and may hence be regulated by Son1p, in addition to Hac1p.

Regulation of Protein Translocation Into and Out of the ER

The observation that UPR-deficient cells expressing CPY*HA exhibit general protein translocation defects revealed another unexpected role for the UPR. As discussed above, our analyses show a delay in the degradation of CPY*HA via ERAD. We consider it likely that these two phenomena are linked as several components of the ER translocon are thought to be both required protein import and export (Wiertz et al., 1996; Pilon et al., 1997; Plemper et al., 1997): UPR-regulated factors shared between import and export may become limiting under conditions as an extra load of an ERAD substrate needs to be attended to.

We confirmed this notion by testing two UPR target genes with known roles in translocation for import and export by increasing their gene dosage. We found that an extra copy of SEC61 was sufficient to alleviate the import defect, but degradation was still compromised (Fig. 7), while an extra copy of KAR2 restored both import and ERAD functions. These results suggest that primarily KAR2 becomes limiting under these conditions. Interestingly, expression of the mouse major histocompatibility complex class I heavy chain H-2Kb in yeast also displayed rapid ER degradation that is dependent on UPR function (Casagrande et al., 2000). In apparent contrast to our results, KAR2 overexpression showed no improvement of heavy chain degradation in Δire1 cells. We also observed, however, that more dramatic overexpression of KAR2 can actually inhibit the degradation of CPY*HA in wild-type cells (Spear, E.D., and D.T.W. Ng, unpublished results; J. Brodsky, personal communication). Taken together, these data suggest that a precise and balanced modulation of UPR targets is required for optimal ERAD activity.

These results serve to illustrate a simple approach that possibly can be more broadly exploited to study pathways under UPR control. Starting with a UPR-deficient strain, one imposes a specific nonlethal stress that in wild-type cells would elicit a moderate UPR induction (in this case, moderate CPY*HA expression). With the UPR off, key components become limiting. As defects are detected, it becomes possible to test the contribution of specific UPR target genes by increasing their gene dosage. Only some of many genes required for any given function are regulated by the UPR (Travers et al., 2000) which, we surmise, are likely to comprise those components that are rate limiting under uninduced conditions. Thus, the approach may allow to identify key regulatory components even for processes that require the contribution of multiple genes.

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