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Amino Acids Induce Peptide Uptake Via Accelerated Degradation of CUP9, the Transcriptional Repressor of the PTR2 Peptide Transporter

Zanxian Xia^{‡†¶}, Glenn C. Turner^{§¶}, Cheol-Sang Hwang[‡], Christopher Byrd[¥],
and Alexander Varshavsky^{‡1}

[¶]These authors contributed equally to this work.

[‡]From Division of Biology, California Institute of Technology, Pasadena, CA 91125; [§]Cold Spring Harbor Laboratory, Cold Spring Harbor, NY 11724; [¥]Wilson, Sonsini, Goodrich & Rosati, 650 Page Mill Road, Palo Alto, CA 94304

[†]Present address: Center for Stem Cell and Regenerative Medicine, University of Southern California, Los Angeles, CA 90033.

¹Address correspondence to: A. Varshavsky, Div. of Biology, California Inst. of Technology, Pasadena, CA 91125. Tel: 626-395-3785; Fax: 626-440-9821; Email: avarsh@caltech.edu

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Multiple pathways link expression of PTR2, the transporter of di- and tripeptides in the yeast *Saccharomyces cerevisiae*, to the availability and quality of nitrogen sources. Previous work has shown that induction of PTR2 by extracellular amino acids requires, in particular, SSY1 and PTR3. SSY1 is structurally similar to amino acid transporters but functions as a sensor of amino acids. PTR3 acts downstream of SSY1. Expression of the PTR2 peptide transporter is induced not only by amino acids but also by dipeptides with destabilizing N-terminal residues. These dipeptides bind to UBR1, the ubiquitin ligase of the N-end rule pathway, and allosterically accelerate the

UBR1-dependent degradation of CUP9, a transcriptional repressor of PTR2. UBR1 targets CUP9 through its internal degron. Here we demonstrate that the repression of PTR2 by CUP9 requires TUP1 and SSN6, the corepressor proteins that form a complex with CUP9. We also show that the induction of PTR2 by amino acids is mediated by the UBR1-dependent acceleration of CUP9 degradation that requires both SSY1 and PTR3. The acceleration of CUP9 degradation is shown to be attained without increasing the activity of the N-end rule pathway toward substrates with destabilizing N-terminal residues. We also found that GAP1, a general amino acid transporter, strongly contributes to the induction of

PTR2 by Trp. While several aspects of this complex circuit remain to be understood, our findings establish new functional links between the amino acids-sensing SPS system, the CUP9-TUP1-SSN6 repressor complex, the PTR2 peptide transporter and the UBR1-dependent N-end rule pathway.

Biological processes addressed by this study include the mechanisms and regulation of peptide import. Peptides can serve as a source of amino acids and nitrogen in all organisms. The import of di- and tripeptides (di/tripeptides) in the yeast *Saccharomyces cerevisiae* has been shown to be regulated by the N-end rule pathway, one proteolytic pathway of the ubiquitin (Ub)-proteasome system (1-4). The N-end rule relates the *in vivo* half-life of a protein to the identity of its N-terminal residue (reviewed in refs. (5-7)). Although prokaryotes lack the Ub system, they still contain the N-end rule pathway, albeit Ub-independent versions of it (6,8). In eukaryotes, this pathway recognizes several kinds of degradation signals (degrons), including a distinct set called N-degrons (5-7). Specific N-degrons that are active (recognized) in a cell give rise to that cell's N-end rule. The determinants of an N-degron in a substrate protein are a destabilizing N-terminal residue that bears the unmodified N-terminal amino group, a substrate's internal Lys residue (the site of formation of a poly-Ub chain), and a conformationally mobile region in a vicinity of this residue (9-11).

The N-end rule has a hierarchic structure. In eukaryotes, N-terminal Asn and Gln are tertiary destabilizing residues in that they function through their enzymatic deamidation, to yield the secondary destabilizing N-terminal residues Asp and Glu (12). Destabilizing activity of N-terminal Asp and Glu requires their conjugation to Arg, one of the primary destabilizing residues, by the *ATE1*-encoded Arg-tRNA-protein transferase (arginyl-transferase or R-transferase) (13-17). In mammals and other eukaryotes that produce nitric oxide (NO), the set of arginylated residues contains not only Asp and Glu but also N-terminal Cys, which is arginylated after its

oxidation to Cys-sulfinate or Cys-sulfonate (13). The *in vivo* oxidation of N-terminal Cys requires NO, as well as oxygen (O₂) or its derivatives (14,17). The N-end rule pathway is thus a sensor of NO, through the ability of this pathway to destroy proteins with N-terminal Cys, at rates controlled by NO, O₂ and their derivatives.

E3 Ub ligases of the N-end rule pathway are called N-recognins (7,18-20). They recognize (bind to) primary destabilizing N-terminal residues. (The term "Ub ligase" denotes either an E2-E3 holoenzyme or its E3 component.) At least four N-recognins, including UBR1, mediate the N-end rule pathway in mammals and other multicellular eukaryotes (7). The known N-recognins share a ~70-residue motif called the UBR box. Mouse UBR1 and UBR2 are the sequelogous (similar in sequence) 200 kDa RING-type E3 Ub ligases that are 47% identical. Several other mammalian N-recognins, either confirmed or putative ones, are HECT-type or SCF-type Ub ligases that share the UBR motif with the RING-type UBR1 and UBR2 but are largely nonsequelogous to them otherwise (21,22). (*A note on terminology*: "sequelog" and "spalog" denote, respectively, a sequence that is similar, to a specified extent, to another sequence, and a three-dimensional structure that is similar, to a specified extent, to another three-dimensional structure (23). Besides their usefulness as separate terms for sequence and spatial similarities, the rigor-conferring advantage of "sequelog" and "spalog" is their *evolutionary neutrality*, in contrast to interpretation-laden terms such as "homolog", "ortholog" and "paralog". The latter terms are compatible with the sequelog/spalog terminology and can be used to convey understanding about functions and common descent, if this (additional) information is available.)

The functions of the N-end rule pathway include: (i) The sensing of heme, owing to inhibition, in both yeast and mammals, of the ATE1 R-transferase by heme (Fe³⁺-heme), which also inhibits N-recognins, the latter at least in yeast. (ii) The sensing of NO and O₂, and the resulting control of signaling by transmembrane receptors, through the conditional, NO/O₂-mediated degradation of G-protein regulators RGS4, RGS5 and RGS16.

(iii) Control of the import of short peptides, through degradation (modulated by peptides) of CUP9, the repressor of import. (iv) Fidelity of chromosome segregation, through degradation of a separase-produced cohesin fragment. (v) Regulation of apoptosis, through degradation of a caspase-processed inhibitor of apoptosis. (vi) A multitude of processes mediated by the transcription factor c-FOS, a conditional substrate of the N-end rule pathway. (vii) Regulation of the human immunodeficiency virus (HIV) replication cycle, through degradation of HIV integrase. (viii) Regulation of meiosis, spermatogenesis, neurogenesis and cardiovascular development in mammals, and leaf senescence in plants (refs. (2-4,13,14,16-19,24-26) and refs. therein). Mutations in human UBR1, one of E3s of the N-end rule pathway, are the cause of Johansson-Blizzard Syndrome (JBS), which comprises mental retardation, physical malformations, and severe pancreatitis (27).

The N-end rule pathway of *S. cerevisiae* is mediated by a single N-recognin, UBR1, a 225 kDa sequelog of mammalian UBR1 and UBR2 (2,28). *S. cerevisiae* UBR1 contains at least three substrate-binding sites. The type-1 site is specific for basic N-terminal residues of polypeptides (Arg, Lys, His). The type-2 site is specific for bulky hydrophobic N-terminal residues (Trp, Phe, Tyr, Leu, Ile). The third binding site of UBR1 recognizes an internal (non-N-terminal) degron in target proteins. The third binding site of UBR1 is autoinhibited but can be allosterically activated through a conformational change that is caused by the binding of short peptides, such as dipeptides, to the other, type-1 and type-2, binding sites of UBR1. The known substrate of the third binding site of UBR1 is CUP9 (1-3), a homeodomain protein and transcriptional regulator, largely a repressor, of more than 30 genes in *S. cerevisiae*¹. The regulon of CUP9 includes *PTR2*, a gene encoding the main transporter of di/tripeptides (29,30). The reversal of UBR1 autoinhibition by imported di/tripeptides with destabilizing N-terminal residues accelerates the UBR1-dependent ubiquitylation of CUP9, leads to its faster degradation, and thereby causes a derepression of *PTR2*. The resulting positive-feedback circuit

allows *S. cerevisiae* to detect the presence of extracellular peptides and to react by increasing their uptake (2,3).

The evolution of peptide-import circuits is under conflicting selective pressures, as the ability of a cell to import peptides confers both a benefit (utilization of peptides as food) and a vulnerability to toxins that resemble short peptides. While *PTR2* is the major transporter of di/tripeptides in *S. cerevisiae*, some di/tripeptides can also be imported (with a low efficacy) by *DAL5*, whose major function is the import of other nitrogen sources, such as allantoin and ureidosuccinate (4). Another set of *S. cerevisiae* peptide transporters comprises *OPT1* (*HGT1*) and *OPT2*, which have partially overlapping functions, do not import di/tripeptides but can import peptides of 4-5 residues. In addition, *OPT1* (*HGT1*) is a high-affinity importer of glutathione, a “noncanonical” tripeptide (ref. (31) and refs. therein). Similarly to the *PTR2* transporter of di/tripeptides, the expression of *OPT2* is downregulated by CUP9, whereas the expression of *OPT1* (*HGT1*) is independent of CUP9 (31). In addition to *PTR2* and *OPT2*, the N-end rule pathway also controls the expression of *DAL5*, but in a manner opposite to that of the other two transporters: whereas CUP9 is a transcriptional repressor of *PTR2* and *OPT2*, CUP9 apparently upregulates the expression of *DAL5* (29,31). It is unknown whether CUP9 downregulates a repressor of *DAL5* or whether CUP9 acts, in the context of *DAL5*, as a transcriptional activator.

The induction of the *PTR2* peptide transporter by di/tripeptides, a process controlled by the UBR1-CUP9 circuit (3), is just one of regulatory inputs that couple *PTR2* expression to the availability and quality of nutrients. For example, *PTR2* expression is downregulated by certain nitrogen sources, including ammonia, but not by other nitrogen sources, such as urea and allantoin (32). The underlying systems, including the N-end rule pathway, ensure that a cell does not waste resources synthesizing large amounts of the *PTR2* transporter in the absence of extracellular peptides, or when a more efficacious nitrogen source, such as ammonia, is present. *PTR2* is also induced by extracellular amino acids, particularly leucine (Leu) or

tryptophan (Trp) (33). This response is likely to be adaptive in natural habitats, as the amino acids that *S. cerevisiae* (a scavenging heterotroph) encounters outside laboratory tend to be the breakdown products of peptides and thus signify a likely presence of di/tripeptides.

Extracellular amino acids regulate PTR2 through the SPS (SSY1-PTR3-SSY5) pathway (34-40). SSY1, an integral membrane protein and a sensor of amino acids, is a sequelog of amino acid transporters but does not function as a transporter (36,37,41), a disposition that recurs with other nutrient sensors as well (37,42). Both the inferred design of SSY1 and experimental evidence suggest that it is the concentration ratio of an amino acid across the plasma membrane, rather than the absolute concentration of extracellular amino acid that determines the signaling output by SSY1 (36). Activated SSY1 induces expression of a regulon that includes the PTR2 peptide transporter and amino acid transporters such as AGP1, BAP2, BAP3, TAT1, TAT2 and GNP1 (38). PTR3 and SSY5 are peripheral membrane proteins associated with SSY1 (41). SSY5 is a protease regulated, in particular, by PTR3. SSY5 can cleave, and thereby activate, the latent (conditionally cytosolic) transcriptional activators STP1 and STP2, leading to their import into the nucleus and the induction of genes that encode, in particular, amino acid transporters (34-37,39,40,43).

In the present work, we show that an extracellular amino acid such as Trp acts via the SPS system to induce the PTR2-mediated import of di/tripeptides through the acceleration of degradation of CUP9, the repressor of import, by the UBR1-dependent N-end rule pathway. The bulk of this effect of Trp on the rate of CUP9 degradation requires both SSY1 and PTR3. At the same time, no significant activation of the N-end rule pathway toward substrates with destabilizing N-terminal residues (i.e., toward substrates with N-degrons) was observed under these conditions, suggesting a differential regulation of three substrate-binding sites of the UBR1 Ub ligase. We also show that the repression of PTR2 by CUP9 requires the global corepressors TUP1 and SSN6, and that GAP1, a general amino acid transporter, strongly contributes to the induction of PTR2 by

Trp. While several aspects of this complex circuit remain to be understood, our findings establish new functional links between the amino acids-sensing SPS system, the CUP9-TUP1-SSN6 repressor complex, the PTR2 peptide transporter and the UBR1-dependent N-end rule pathway.

EXPERIMENTAL PROCEDURES

Yeast Strains, Plasmids, and Genetic Techniques – The *S. cerevisiae* strains used in this study are described in Table 1. AVY24 and AVY25 were constructed in the background of strain RJD347 (*MAT α ura3-52*; a gift from Dr. R. Deshaies, California Institute of Technology). A PCR-based deletion strategy (44) was employed, using DNA fragments encoding the *myc*₃-URA3-*myc*₃ cassette to precisely delete *SSY1* and *PTR3* open reading frames (ORFs) (*myc*₃ denotes 3 repeats of a nucleotide sequence encoding the *myc* epitope, flanked on either side by 50 bp of sequences identical to the sequences of either the *SSY1* or the *PTR3* gene, respectively). This technique was employed to produce deletions of the *SSY1* and *PTR3* ORFs. We then selected for recombination within the *myc*₃-URA3-*myc*₃ module using 5-fluoroorotic acid (FOA), to produce strains AVY27 (*MAT α ura3-52 ssy1 Δ ::myc₃*) and AVY28 (*MAT α ura3-52 ptr3 Δ ::myc₃*), and verified the identity of these strains by Southern hybridization (data not shown). AVY26 (*MAT α ura3-52 ubr1 Δ ::HisG*) was also a derivative of RJD347 (Table 1). It was constructed using the same PCR-based deletion strategy (44), except a *HisG-URA3-HisG* cassette (see below) was used. Strains AVY30 (*MAT α leu2-3,112 ubr1 Δ ::LEU2*), AVY31 (*MAT α leu2-3,112 cup9 Δ ::LEU2*), and AVY32 (*MAT α LEU2*) (Table 1) were derivatives of RJD350 (*MAT α leu2-3,112*) (a gift from Dr. R. Deshaies).

PCR with *S. cerevisiae* genomic DNA was used to produce a fragment containing *SSN6* flanked by 901 bp and 400 bp of upstream and downstream sequences, respectively. The fragment was ligated to *NotI/EcoRI*-cut pRS314 (45). The same procedure was used to construct a fragment containing *TUP1* flanked by 459 bp and 400 bp of upstream and downstream sequences, respectively, followed by ligation to *SpeI/Clal*-cut pRS416 (45). pSsn6 Δ , which

carried *ssn6Δ::HisG-URA3-HisG*, was constructed by subcloning the *SSN6*-containing *NotI-EcoRI* fragment into *NotI/EcoRI*-cut Bluescript (Stratagene), producing pBlueSSN6. Using PCR, a deletion (from 50 bp upstream of the *SSN6* start codon to position 2,585 of its ORF) was introduced, while simultaneously creating an *EclXI* site at the upstream breakpoint of the deletion. The *EclXI-PstI* fragment of pAS315 (a gift from Dr. A. Sil, University of California, San Francisco) containing *HisG-URA3-HisG*, was then inserted. *In vivo* integration of the *NotI-EcoRI* fragment of the resulting plasmid (pSsn6Δ), carrying *ssn6Δ::HisG-URA3-HisG*, produced a deletion of *SSN6* that spanned the region from -50 to 2,585, relative to start codon. pTup1Δ, which carried *tup1Δ::HisG-URA3-HisG*, was constructed similarly, using the *SpeI-ClaI* fragment containing *TUP1* (from pTUP1), its subcloning into pBluescript (thus yielding pBlueTUP1), and PCR to produce a deletion allele of *TUP1* (from 7 bp upstream of its start codon to position 2,016 of the *TUP1* ORF) that contained a (PCR-added) *BamHI* site at the deletion breakpoint. The *BamHI-EcoRI* fragment of pAS135 was then inserted into pBlueTUP1. *In vivo* integration of the *SpeI-XhoI* fragment of the resulting plasmid (carrying *tup1Δ::HisG-URA3-HisG*) produced a deletion of *TUP1*, from -7 to +2,016, relative to start codon. To select for recombination/excision within the integrated *HisG-URA3-HisG* cassette and thereby to produce *ssn6Δ::HisG* and *tup1Δ::HisG* alleles, 5-fluoroorotic acid (FOA)-resistant yeast colonies were isolated, and the identities of strains were verified by Southern hybridization (data not shown). This procedure was performed in the JD52 background to construct strains AVY60 and AVY61; in the JD55 (*ubr1Δ*) background (Table 1) (24) to construct AVY62 and AVY63; in the AVY50 background (1) to construct AVY64 and AVY65; and in the AVY51 background (1) to construct AVY66 and AVY67 (Table 1).

SSN6 tagged at the C-terminus with two copies of the myc epitope (*SSN6_{myc2}*) and expressed from its own promoter on a high copy (2μ) plasmid was a gift from Dr. R. Zitomer (State University of New York, Albany). For plasmid-based complementation assays,

plasmids encoding SSN6 (pSSN6) and TUP1 (pTUP1) were constructed using standard techniques. ^fDHFR-Ub^{K48R}-CUP9_{NSF} was expressed from the P_{MET25} promoter in the plasmid pMET416UPRCUP9_{NSF}, based on the low copy (*CEN*) vector p416MET25 (3). The previously described (28) pUb23-X plasmids expressed X-βgal reporter proteins (derived from the corresponding Ub-X-β-galactosidase fusions; X=Met, Ala, His or Tyr) from the P_{GALI} promoter in a high copy vector.

Coimmunoprecipitation, Pulse-Chase, β-Galactosidase and Northern Hybridization Assays – For CUP9-SSN6 coimmunoprecipitation assays, *S. cerevisiae* carrying the indicated plasmids were grown to A₆₀₀ of ~1 in SD medium containing auxotrophic supplements. Cells from a 10-ml culture were harvested by centrifugation, washed in 1 ml of water, and resuspended in 0.8 ml of ice-cold buffer C (0.25 M NaCl, 1 mM EDTA, 50 mM HEPES, pH 7.5) containing ovalbumin at 1 mg/ml. Cells were disrupted by vortexing with 0.5 ml (packed volume) of 0.5-mm glass beads in the presence of protease inhibitors (Roche). Extracts were prepared and immunoprecipitations were carried out as described (46), with two modifications. First, after incubation of an extract with anti-flag beads (Sigma, St. Louis, MO) for 45 min at 4°C, the resulting immunoprecipitate was washed once with 1 ml of buffer C, and then twice more with 1 ml of buffer C lacking ovalbumin. Second, immunoprecipitates were eluted from anti-flag beads through a 15-min incubation, at room temperature, with the flag peptide at 0.5 mg/ml, in buffer C lacking ovalbumin. An equal volume of 2 x SDS-PAGE sample buffer was added, followed by SDS-13% PAGE and immunoblotting with anti-myc antibody (Covance, Berkeley, CA), followed by a secondary horseradish peroxidase-conjugated goat anti-mouse antibody (BioRad, Hercules, CA) at 1:1,000 dilution, and thereafter a detection of the latter antibody using ECL assay (GE Healthcare, Piscataway, NJ).

For pulse-chase assays, *S. cerevisiae* was grown to A₆₀₀ of ~0.8 in SHM medium (2% glucose, 0.1% allantoin, 0.17% yeast nitrogen base without amino acids and without ammonium sulfate) that either contained or

lacked tryptophan (Trp) (98 μ M; 20 μ g/ml), an inducing amino acid (33). Cells were harvested, washed in 0.8 ml of SHM \pm Trp, resuspended in 0.4 ml of the same medium, and labeled for 5 min at 30°C with 0.16 mCi of 35 S-EXPRESS (Perkin-Elmer, Waltham, MA, USA). Thereafter cells were pelleted, resuspended in fresh SHM \pm tryptophan containing 4 mM L-methionine and 2 mM L-cysteine, and incubated further at 30°C. Samples of 0.1 ml were taken at indicated time points and transferred to chilled tubes, each containing 0.5 ml of 0.5-mm glass beads, 0.7 ml of ice-cold lysis buffer (1% Triton-X100, 0.15 M NaCl, 5 mM EDTA, 50 mM HEPES, pH 7.5), and a mixture of protease inhibitors (final concentrations: 1 mM freshly prepared phenylmethylsulfonyl fluoride, 2 μ g/ml aprotinin, 0.5 μ g/ml leupeptin, and 0.7 μ g/ml pepstatin). Extracts were prepared and immunoprecipitations carried out as described (46), using anti-flag M2 beads (Sigma). Immunoprecipitated proteins were fractionated by SDS-13% PAGE, followed by autoradiography. Some of pulse-chase assays were carried out in the presence of cycloheximide (at 0.2 mg/ml) during the chase.

For β -galactosidase (β gal) activity assays, *S. cerevisiae* carrying a pUB23-X plasmid (28) and expressing, from the P_{GALI} promoter, an X- β gal test protein (derived from Ub-X- β -galactosidase, with X=Met, Ala, His or Tyr) (47) were grown in SHM medium overnight. Cultures were diluted to A_{600} of \sim 0.1 in SHM containing 2% galactose instead of glucose and either containing or lacking 98 μ M Trp, followed by growth until A_{600} of \sim 0.5. Cell extracts were thereafter prepared and processed for assays with o-nitrophenyl o-nitrophenyl- β -D-galactopyranoside (ONPG) as previously described (28). For Northern analyses of *PTR2* induction by amino acids, cells were grown to A_{600} of \sim 0.6 in SHM containing or lacking 98 μ M (20 μ g/ml) Trp. Total RNA was prepared as described previously (15), using RNeasy Mini Kit (Qiagen, Valencia, CA). Samples containing 25 μ g of RNA were electrophoresed in 1% formaldehyde-agarose gels, followed by blotting for Northern analysis (3). Northern assays of the effects of deletions of specific genes on *PTR2* expression were carried out identically, in the absence of added Trp.

RESULTS AND DISCUSSION

Repression of PTR2 Transcription by CUP9 Involves the CUP9-TUP1-SSN6 Corepressor Complex – UBR1, the E3 Ub ligase of the *S. cerevisiae* N-end rule pathway, regulates the import of peptides by controlling, through degradation of the transcriptional repressor CUP9, the expression of the peptide transporter *PTR2* (1). CUP9 was identified using a bypass screen for mutations that restored dipeptide uptake and *PTR2* expression in *ubr1* Δ cells (1). Further analysis, described below, of our collection of bypass mutants identified additional isolates that did not belong to the *CUP9* complementation group. Many of these non-*cup9* mutants (Fig. 1A) exhibited flocculation (aggregation during growth in liquid culture) and had a reduced mating ability. These phenotypes are characteristic of a defect in either *TUP1* or *SSN6*, which encode two “global” corepressors that form a complex interfering with transcription initiation (48-50)). The TUP1-SSN6 complex, which does not bind to DNA by itself, is recruited to individual promoters through interactions with specific DNA-binding proteins. One such protein is the mating-type regulator MAT α 2 (51,52). In the absence of either TUP1 or SSN6, the MAT α 2-mediated repression of *MAT α* -specific genes is defective, resulting in mating-defective, pseudodiploid cells (48).

To determine whether the above non-*cup9* isolates (Fig. 1A) were either *tup1* or *ssn6* mutants, they were transformed with low-copy plasmids that expressed either *TUP1* or *SSN6* from their natural promoters. These plasmids indeed suppressed the clumpy-growth and peptide-uptake phenotypes of all of the flocculating bypass mutants. The entire collection of our 201 bypass mutants (including *cup9* mutants) was partitioned into complementation groups using a combination of the above plasmid-based complementation tests as well as mating-based complementation assays (the latter could not be comprehensive, owing to mating deficiency of some isolates). The 201 bypass suppressors were found to belong to 3 complementation groups, defined by *CUP9*, *TUP1* and *SSN6*. Apart from genes required for cell viability, it is likely that our

(nonconditional) screen was an exhaustive one, given the numbers of isolates in each of the 3 complementation groups thus identified: 104, 67, and 30 isolates of *cup9*, *ssn6* and *tup1* mutants, respectively (Fig. 1A).

To verify the role of *TUP1* and *SSN6* in regulating peptide uptake, we tested the effects of disrupting either one of these genes on the ability of a *ubr1Δ* strain to import dipeptides. Congenic [*tup1Δ ubr1Δ*] and [*ssn6Δ ubr1Δ*] double mutants were constructed using homologous recombination in the lysine (Lys)-auxotrophic (*lys2*) genetic background, and the identities of mutants were verified by Southern hybridization (data not shown). The ability of these Lys-requiring double mutants to import dipeptides was then assayed by monitoring their growth on a minimal medium containing the dipeptide Lys-Ala as the sole source of Lys (Fig. 1B). A *lys2* strain containing the intact *UBR1* gene was able to import sufficient amounts of Lys-Ala dipeptide to support its growth on this medium, whereas a congenic [*ubr1Δ lys2*] strain could not grow under these conditions (it could be rescued by free Lys) (Fig. 1B). In contrast to the [*ubr1Δ lys2*] strain, both [*tup1Δ ubr1Δ lys2*] and [*ssn6Δ ubr1Δ lys2*] triple mutants formed colonies on Lys-Ala plates, as did a [*cup9Δ ubr1Δ lys2*] strain (Fig. 1B). Thus, a deletion of either *CUP9*, or *TUP1*, or *SSN6* is each capable of restoring dipeptide uptake to a *ubr1Δ* strain. The slightly smaller colonies formed by the [*ssn6Δ ubr1Δ lys2*] strain on Lys-Ala plates could be caused by a growth defect of *ssn6Δ* strains (49), a phenotype that we also observed with the [*ssn6Δ ubr1Δ lys2*] strain on a control (Lys-supplemented) plate (Fig. 1B).

The fact that *cup9*, *tup1* and *ssn6* mutants all bypassed the block to dipeptide uptake in a *ubr1Δ* strain suggested that *CUP9* represses *PTR2* expression by functioning as a part of a *TUP1/SSN6*-containing complex. To address this model, we compared the levels of *PTR2* mRNA in *ssn6Δ* and *tup1Δ* mutants with those in other strains, in the absence of added *PTR2* inducers such as dipeptides or amino acids. In a strain retaining all 5 of the studied genes, *PTR2*, *UBR1*, *CUP9*, *SSN6* and *TUP1*, and also in a *ubr1Δ* strain, the level of *PTR2* mRNA was low enough to be undetectable at the

sensitivity of Northern assays used (Fig. 2A, lanes a, b; compare with lanes c-i). With *ubr1Δ* strains, no *PTR2* mRNA could be detected even by higher-sensitivity Northern (data not shown), whereas wild-type (*UBR1*) strains, in the absence of either added dipeptides or “inducing” amino acids, contained low but detectable levels of *PTR2* mRNA, as shown previously (1,3). In contrast, the levels of *PTR2* mRNA were high in the absence of either *CUP9*, or *SSN6*, or *TUP1*, irrespective of the presence or absence of *UBR1* (Figs. 2A, lanes c-i; compare with lanes a, b; see also Fig. 3A, lane c versus lane i). Thus, the tight block to dipeptide uptake in a *ubr1Δ* strain, caused by increased levels of the *CUP9* repressor, could be bypassed not only through the removal of *CUP9*, but also through the removal of either *SSN6* or *TUP1* (Fig. 2A).

High levels of *PTR2* mRNA in the absence of either *CUP9*, *SSN6*, or *TUP1* (Figs. 2A and 3A) indicated that each of these proteins was essential for repression of the *PTR2* gene. Both the demonstrated functional dependence of *CUP9* on *SSN6/TUP1* (Figs. 1 and 2A) and the close analogy between the homeodomain repressors *CUP9* and *MATα2*, the latter of which is known to interact with *SSN6/TUP1* (see above), suggested that *CUP9* also interacted with *SSN6/TUP1*. To address this experimentally, we carried out a coimmunoprecipitation assay, using epitope-tagged *SSN6* and *CUP9* (Fig. 2B). Since overexpression of wild-type *CUP9* inhibits cell growth, most likely owing to its interactions with DNA (data not shown), we employed a previously constructed nontoxic derivative of *CUP9* (3). *CUP9* interacts with DNA through its homeodomain motif. Based on the previously studied interactions of other homeodomain proteins with DNA, an Asn (N) → Ser (S) substitution at position 265, within the recognition helix of the *CUP9* homeodomain, was predicted to strongly reduce the affinity of *CUP9* for DNA without causing a significant structural perturbation (51). This nontoxic *CUP9* derivative, tagged at the C-terminus with the flag epitope, was termed *CUP9_{NSF}* (3). Extracts were prepared from *S. cerevisiae* stably expressing both *CUP9_{NSF}* and *SSN6* (*SSN6_{myc2}*) (the latter C-terminally

tagged with a double myc epitope), followed by immunoprecipitation with anti-flag antibody, SDS-PAGE, and immunoblotting with anti-myc antibody. *SSN6_{myc2}* was coimmunoprecipitated with *CUP9_{NSF}* in this assay (Fig. 2B, lane c). No *SSN6_{myc2}* was detected in immunoprecipitates of extracts from cells that did not also express *CUP9_{NSF}* (Fig. 2B, lane b), and the expression levels of *CUP9_{NSF}* and *SSN6_{myc2}* were similar in cells that coexpressed the two proteins (data not shown). Thus, in agreement with the above expectation, these results (Fig. 2B) suggested that untagged *SSN6* and untagged *CUP9*, at their (lower) natural expression levels in *S. cerevisiae*, would also form a complex that is analogous to the *SSN6/MATα2* complex (50), a model that accounts for the observed functional interdependence among *CUP9*, *SSN6* and *TUP1* (Figs. 1B, 2A and 3A). While we did not examine the immunoprecipitated *CUP9_{NSF}*-containing complexes for the presence of *TUP1*, extensive previous work has established that *SSN6* and *TUP1* form a stable complex both *in vivo* and as purified proteins (49), indicating that it is the complex of *SSN6* and *TUP1* that interacts with *CUP9*, similarly to the previously characterized interaction between *SSN6/TUP1* and *MATα2*, another homeodomain repressor (50).

Both CUP9 and UBR1 Are Required for Inducibility of PTR2 by Amino Acids – When an amino acid, particularly a bulky hydrophobic one such as Leu or Trp, is added to a culture of *S. cerevisiae* growing on a poor nitrogen source such as, for example, allantoin, cells induce their *PTR2* gene and thus increase their capacity for the import of di/tripeptides (33). Effective concentrations of extracellular Leu or Trp can be as low as 1 μM (33). Previous work has shown that an accelerated degradation of *CUP9* by the *UBR1*-dependent N-end rule pathway in response to imported dipeptides with destabilizing N-terminal residues, strongly enhances *PTR2* expression (3). This result led us to consider whether other signals that up-regulate *PTR2* expression, e.g., amino acids, may also act through acceleration of *CUP9* degradation.

To begin addressing this possibility, we examined the effects of Trp, a strongly inducing amino acid, on *PTR2* mRNA levels in several *S. cerevisiae* strains, *AVY32*, *AVY24*, *AVY25*, *AVY30* and *AVY31*, that were constructed to be completely prototrophic, and in particular did not require extracellular amino acids for viability (Table 1). These strains were grown in allantoin-based media (SHM) containing or lacking Trp. Allantoin is a nonrepressing nitrogen source (33), so the expression of *PTR2* was not influenced by the nitrogen catabolite repression under these conditions (53). *PTR2* mRNA levels were extremely low in wild-type [*UBR1 CUP9*] cells grown in medium lacking amino acids, but in the presence of 98 μM (20 μg/ml) Trp the expression of *PTR2* was greatly increased (Fig. 3A, lanes a, b). By contrast, in the absence of *CUP9* (in a *cup9Δ* strain), *PTR2* was highly expressed in either the presence or absence of Trp (Fig. 3A, lanes i, j; compare with lanes a, b). Finally, *PTR2* mRNA was undetectable in a *ubr1Δ* strain grown in either the presence or absence of Trp (Fig. 3A, lane c, d). Thus, both *UBR1* and *CUP9* are required for inducibility of *PTR2* expression by an amino acid.

Three genes, *SSY1*, *PTR3*, and *SSY5*, are a part of the SPS amino acid-sensing pathway in *S. cerevisiae* (see Introduction). We examined the influence of added Trp on the levels of *PTR2* mRNA in *ssy1Δ* and *ptr3Δ* strains, comparing the consequences of deleting these genes with the effects of deleting either *CUP9* or *UBR1*. The induction of *PTR2* by Trp was dramatically reduced in both *ssy1Δ* and *ptr3Δ* strains (Fig. 3A, lanes e-h). However, a weak Trp-mediated increase in *PTR2* expression was still observed in these strains, in contrast to the complete absence of *PTR2* expression (and of Trp effect) in *ubr1Δ* cells, and in contrast to a high level of *PTR2* mRNA in *cup9Δ* cells (Fig. 3A). Thus the bulk (but apparently not all) of the Trp effect on the expression of *PTR2* requires *SSY1* and *PTR3*.

We next examined whether the general amino acid permease *GAP1* (37,54) was required for Trp-mediated induction of *PTR2* mRNA. Expression of *PTR2* in wild-type cells and a congenic *gap1Δ* strain was measured by Northern analysis as a function of time after the addition of Trp. Wild-type cells exhibited a

strong induction of *PTR2* by 20 min in the presence of Trp (Fig. 3D, E, lanes a-c). Remarkably, the induction of *PTR2* by Trp, while still occurring in the absence of GAP1, was found to be strongly decreased (Fig. 3D, E, lanes d-f; compare with lanes a-c). To the best of our knowledge, this is the first indication of a major involvement of GAP1 in the amino acid-mediated induction of the *PTR2* peptide transporter. Being an inducer of *PTR2* and other transporter genes via the amino acid sensor SSY1, the extracellular Trp exerts its effect without its import across the plasma membrane (see Introduction). Therefore the absence of GAP1 as an importer of Trp would not be expected, *a priori*, to influence the *PTR2*-inducing effect of Trp. Moreover, Trp can also be imported by the Trp-specific permease TAP2 (SCM2) (55). In sum, the observed decrease (but not elimination) of *PTR2* induction in the absence of *GAP1* (Fig. 3D, E, lanes a-f) stems from a function of GAP1 that is separate from its role as a transporter of amino acids. Indeed, *S. cerevisiae* GAP1 is known to function as an amino acid sensor for activation of protein kinase-A (PKA) targets. The PKA and SCH9 kinases have both overlapping and distinct functions in controlling the adaptation of *S. cerevisiae* to nutrient availability, and GAP1 plays a role at least in the PKA-mediated part of this circuit (54,56). Thus, while the mechanistic understanding of GAP1's involvement in the induction of *PTR2* by Trp remains to be attained, the above finding (Fig. 3D, E) revealed a new aspect of *PTR2* control and yet another function of GAP1 as a nutritional regulator.

We also measured, in the *gap1* background, the influence of STP1 and/or STP2 on the Trp-mediated induction of *PTR2* mRNA. STP1 and STP2 are two sequelogous, conditionally active transcriptional activators that mediate the signaling by the SPS pathway (see Introduction) (35,39,40,43). Northern analyses of *PTR2* expression as a function of time after addition of Trp showed that the Trp-mediated induction of *PTR2* was largely retained in a [*gap1*Δ *stp1*Δ] strain (which lacked the STP1 transcriptional activator), in comparison to congenic *gap1*Δ strain, whereas the STP2 transcriptional activator was essential for *PTR2* induction by Trp: both a [*gap1*Δ *stp2*Δ] mutant

and a triple mutant [*gap1*Δ *stp1*Δ *stp2*Δ] did not exhibit a significant induction of *PTR2* by Trp, in contrast to parental *gap1*Δ and [*gap1*Δ *stp1*Δ] strains (Fig. 3D, E). Thus, as could be expected from the presence of STP1/2-recognized UAS_{AA} nucleotide sequence motifs (57) upstream of the *PTR2* gene (data not shown), the induction of *PTR2* proceeds both through its transcriptional upregulation, in particular by the Trp-activated STP2 (Fig. 3D, E), and through a decrease in its repression, owing to the Trp-induced acceleration of degradation of the CUP9 repressor by the N-end rule pathway, as described below (Fig. 4).

The SPS Pathway Induces PTR2 Expression Through Acceleration of CUP9 Degradation – Some of the above results (Fig. 3A) suggested that the induction of *PTR2* transporter by amino acids might occur through an increase in the rate of degradation of CUP9 repressor. A precedent for this possibility was the previously demonstrated acceleration of CUP9 degradation by dipeptides with destabilizing N-terminal residues (3). In the latter case, the bulk of CUP9 degradation, both before and after the addition of dipeptides to the medium, was carried out by the UBR1-dependent N-end rule pathway (3). To determine whether the rate of CUP9 degradation was altered by a *PTR2*-inducing amino acid such as Trp, we carried out pulse-chase assays with CUP9. These experiments utilized CUP9_{NSF}, the above-described nontoxic, single-residue mutant of CUP9. CUP9_{NSF} was expressed as a ¹DHFR-Ub^{K48R}-CUP9_{NSF} fusion, where ¹DHFR was the N-terminally flag-tagged mouse dihydrofolate reductase. Deubiquitylating enzymes (DUBs) cotranslationally cleave this fusion at the Ub^{K48R}-CUP9_{NSF} junction, yielding the long-lived reference protein ¹DHFR-Ub^{K48R} and the test protein CUP9_{NSF} (3). The reference ¹DHFR-Ub^{K48R} served as a “built-in” internal control for variations in expression levels and immunoprecipitation efficiency. This generally applicable method, called the UPR (Ub-protein-reference) technique (refs. (10,47) and refs. therein), increases the accuracy of pulse-chase assays. The *in vivo* degradation of CUP9_{NSF} was indistinguishable from that of wild-type CUP9, and was also not altered significantly by the incorporation of this protein into the above

(cotranslationally processed) UPR-type fusion (3).

We expressed CUP9_{NSF} from a *URA3*-bearing plasmid in *S. cerevisiae* strains that did not require extracellular amino acids for viability (RJD347, AVY26, AVY27 and AVY28; see Table 1). In wild-type (*UBR1*) cells grown in the SHM medium lacking amino acids, CUP9_{NSF} was degraded with $t_{1/2}$ of ~10 min (Fig. 4A, B, E, F). Remarkably, the addition of Trp (to 98 μ M, or 20 μ g/ml) to the medium resulted in a strong acceleration of CUP9_{NSF} degradation, with $t_{1/2}$ of ~2 min between 0 and 5 min of chase, and even less than 2 min between 5 and 10 min of chase (Fig. 4A, B, E, F). This effect of Trp on the rate of CUP9_{NSF} degradation was reproducible in separate and independent pulse-chase assays, including a set of assays in which cycloheximide, a translation inhibitor, was present during chase (Fig. 4C, G). (The other assays, in Fig. 4A, B, E, F, did not involve cycloheximide.) Although in cycloheximide-based assays (Fig. 4C, G) the CUP9_{NSF} protein was degraded more slowly ($t_{1/2}$ of ~20 min and 3-4 min in the absence and presence of Trp, respectively) than in cycloheximide-free assays (Fig. 4A, B, E, F), the effect of Trp was comparably strong in both cases (Fig. 4). In addition to increasing the accuracy of pulse-chase assays, the UPR technique, specifically its “built-in”, long-lived reference protein, makes it possible to determine the *relative* level of a test protein (measured as the ratio of ³⁵S in a test versus reference protein) at the beginning of chase, i.e., at the end of pulse labeling (10). In Fig. 4E-G, 100% was assigned, in each of three panels, to the relative amount of ³⁵S (relative to the reference protein) of CUP9_{NSF} at time “0” (the end of pulse labeling) in the absence of Trp in the medium. Thus, a value below 100% at time “0” reflects the relative extent of CUP9_{NSF} degradation during the 5-min pulse. In all of pulse-chase assays (Fig. 4E-G), these initial (time-zero) levels of CUP9_{NSF} were found to correlate, consistently, with Trp-induced changes in the rates of CUP9_{NSF} degradation that were measured during the chase.

The presence of *SSY1* (Fig. 4A, E) and also, independently, of *PTR3* (4B, F) was found to be strongly but partially required for the Trp-

accelerated degradation of CUP9_{NSF}.

Specifically, the bulk of enhanced degradation of CUP9_{NSF} upon the addition of Trp was absent in *ssy1* Δ and *ptr3* Δ mutants (Fig. 4A, B, E, F). Interestingly, however, the degradation of CUP9_{NSF} was still detectably accelerated by Trp in these mutant strains (Fig. 4A, B, E, F), in agreement with the weak but still detectable induction, by Trp, of *PTR2* mRNA in *ssy1* Δ and *ptr3* Δ mutants (Fig. 3A). As we observed previously as well (3), CUP9_{NSF} was greatly but not completely stabilized in *ubr1* Δ cells (Fig. 4G and data not shown). Interestingly, the addition of Trp to *ubr1* Δ cells resulted in a weak but still detectable Trp-induced destabilization of the (now long-lived) CUP9_{NSF}, despite the absence of the N-end rule pathway (Fig. 4G, open circles versus open triangles). The “residual” (*UBR1*-independent) instability of CUP9_{NSF} and the “residual” sensitivity of this degradation to Trp suggest that CUP9 may also be targeted, at a much lower rate, by a *UBR1*-independent proteolytic pathway. Our attempts to identify a relevant E3 Ub ligase, through the testing of non-*UBR1* E3 mutants, have not been successful, thus far (data not shown).

The Levels of UBR1 mRNA Are Not Changed Significantly by a PTR2-Inducing Amino Acid – Previous work has shown that the *in vivo* concentration of *UBR1* is rate-limiting for degradation of N-end rule substrates by the *S. cerevisiae* N-end rule pathway (reviewed in ref. (5)). To determine whether the Trp-induced acceleration of CUP9 degradation (Fig. 4) was accompanied by an increased level of the *UBR1* Ub ligase, we used Northern hybridization to compare the levels of *UBR1* mRNA before and after the addition of Trp. Extracellular Trp did not cause a significant alteration in the levels of *UBR1* mRNA (Fig. 3B, C). Together with findings described in the next section (Fig. 5), these results (Fig. 3B, C) strongly suggested that the observed acceleration of CUP9 degradation in the presence of extracellular Trp (Fig. 4) was not caused by increased levels of *UBR1*. In addition, the levels of *UBR1* mRNA remained the same or nearly the same in the absence of CUP9 (Fig. 3B, C), indicating that *UBR1* is not a part of the CUP9 regulon.

Degradation of Reporter Proteins Bearing N-Degrans Is Not Changed

