Heavy metal sensitivities of gene deletion strains for *ITT1* and *RPS1A* connect their activities to the expression of *URE2*, a key gene involved in metal detoxification in yeast

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Abstract

Heavy metal and metalloid contaminations are among the most concerning types of pollutants in the environment. Consequently, it is important to investigate the molecular mechanisms of cellular responses and detoxification pathways for these compounds in living organisms. To date, a number of genes have been linked to the detoxification process. The expression of these genes can be controlled at both transcriptional and translational levels. In baker’s yeast, *Saccharomyces cerevisiae*, resistance to a wide range of toxic metals is regulated by glutathione S-transferases. Yeast *URE2* encodes for a protein that has glutathione peroxidase activity and is homologous to mammalian glutathione S-transferases. The *URE2* expression is critical to cell survival under heavy metal stress. Here, we report on the finding of two genes, *ITT1*, an inhibitor of translation termination, and *RPS1A*, a small ribosomal protein, that when deleted yeast cells exhibit similar metal sensitivity phenotypes to gene deletion strain for *URE2*. Neither of these genes were previously linked to metal toxicity. Our gene expression analysis illustrates that these two genes affect *URE2* mRNA expression at the level of translation.

Introduction

Heavy metals and metalloids comprise a group of elements that are loosely defined as relatively high-density transition metals and metalloids [1], [2]. Different metals are found in varied
concentrations across the environment. Some of these heavy elements, such as iron (Fe), cobalt (Co) and zinc (Zn), are essential nutrients, while others are relatively harmless at low concentrations such as rubidium (Rb), silver (Ag) and indium (In). At higher concentrations, all metals and metalloids derived from natural environment [3] or anthropogenic sources such as phosphate fertilizers, disinfectants, fungicides, sewage sludge, industrial waste, bad watering practices in agricultural lands, and dust from smelters [4], [5] are toxic to living cells [6], [7], [8]. Among these, arsenic (As) is one of the most toxic despite being the twentieth most abundant element on our planet. Its inorganic oxyanion forms including arsenite (As(III)) and arsenate (As(V)) are highly lethal to living organisms [9].

Over the course of evolution, many organisms have found ways, for example, by evolving molecular pathways to survive increased concentrations of metallic toxins in their environment e.g. [10], [11], [12], [13]. Microbes with extreme adaptation to heavy metals use detoxification pathways to reduce toxic metals to a lower redox state, which lessens their mobility and toxicity [14]. The baker’s yeast, *Saccharomyces cerevisiae* possesses an effective mechanism to negate heavy metal and metalloids toxicity, allowing it to survive a broad range of toxic stress scenarios [15], [16]. This makes yeast an ideal model organism to study molecular mechanisms of the stress response that drive detoxification processes.

The glutathione S-transferases (GSTs) are key enzymes that mediate the resistance of *S. cerevisiae* to a wide range of heavy metals and metalloids. Yeast *Ureidosuccinate Transport 2* (*URE2*) gene product is structurally homologous to mammalian GST and is a major player in the detoxification of *S. cerevisiae* against toxic metals through its glutathione peroxidase activity [17], [18]. For detoxification purposes, GST proteins catalyze the conjugation of the reduced form of glutathione (GSH) to xenobiotic substrates [19]. The deletion strain for *URE2* is hypersensitive to a wide range of heavy metals and metalloids including As, Cd and Ni [19], [20]. In this report, we show that the deletion of either *ITT1* (inhibitor of translation termination 1) or *RPS1A* (small ribosomal subunit protein 10), makes the cells more sensitive to As(III), cadmium (Cd) and nickel (Ni) [19], [20]. In this report, we show that the deletion of either *ITT1* (inhibitor of translation termination 1) or *RPS1A* (small ribosomal subunit protein 10), makes the cells more sensitive to As(III), cadmium (Cd) and Ni suggesting a functional connection of these two genes with heavy metal toxicity. *Itt1p* is known to modulate the efficiency of translation termination through physical interactions with two eukaryotic release factors eRF1 (Sup45p) and eRF3 (Sup35p) [21]. *Rps1Ap* is a constituent of the small ribosomal subunit; little information is known about its function in *S. cerevisiae* [22]. Neither of these genes had previously been linked to heavy metal toxicity. Overall, we provide evidence that the connection for *ITT1* and *RPS1A* with heavy metal toxicity is through their influence on the translation of *URE2* gene.

**Materials and methods**

**Strains and plasmids used in this study**

Yeast, *S. cerevisiae*, mating type (a) MATα strain Y4741 (*MATα orf3::KanMAX4 his3Δ1 leu2Δ0 met15Δ0 ura3Δ0*) and mating type (α) MATα strain, BY7092 (*MATα can1Δ::STE2pr-Sp_his5 lyp1Δ his3Δ1 leu21Δ0 ura3Δ0 met15Δ0*) [23] were utilized for this study. The yeast MATα knockout (YKO) collection [23] and PCR-based transformed cells were used as a source of gene deletion mutants; the open reading frame (ORF) collection [24] was used for over expression plasmid vectors. Yeast GFP Clones [25] were modified for Western blot analysis. pAG25 plasmid containing the Nourseothricin Sulfate (clonNAT) resistance gene was used as a DNA template in PCR to generate gene knockouts. *Escherichia coli* strain DH5α was used to replicate different plasmids [26]. Two plasmids were used carrying the β-galactosidase open reading frame for quantification of *URE2*-IRES and cap-dependent translation activities. p281-4-*URE2* contained a *URE2*-IRES region which was fused with *LacZ* (β-galactosidase) gene from *E. coli*.
and p281 contained only the β-galactosidase gene as a control for cap-dependent translation [27]. All plasmids carried an ampicillin resistance gene which was used as selectable marker in E. coli and the URA3 marker gene which was used for selection in yeast.

**Media and miniprep**

YP (1% Yeast extract, 2% Peptone) or SC (Synthetic Complete) with selective amino acids (0.67% Yeast nitrogen base w/o amino acids, 0.2% Dropout mix,) either with 2% dextrose or 2% galactose as a source of carbohydrates was used as culture medium for yeast and LB (Lysogeny Broth) was used for E. coli cultures. 2% agar was used for all solid media. Yeast cells were grown at 30°C unless otherwise indicated in the SGA procedure. E. coli cells were grown at 37°C. Yeast plasmid extraction was performed by using yeast plasmid mini kit (Omega Bio-Tek®) and E. coli plasmid extraction was carried out by using GeneJET plasmid miniprep kit (Thermofisher®) according to the manufacturer’s instructions.

**Gene knockout and DNA transformation**

Mutant strains were either selected from the library of gene deletions [23] in MATa haploid form or a PCR-based gene knockout approach was used to achieve gene deletion. Targeted gene knockout strains were generated by PCR-based gene deletion strategy utilizing the clonNAT selection gene [28], [29]. Plasmid and gene transformation were performed by using a chemical-transformation strategy (LiOAc method) and confirmed via colony PCR [30], [31].

**Chemical sensitivity**

Colony count assay (spot test) was performed to estimate the cell sensitivities, based on their ability to give rise to colonies. Strains were grown in liquid YPD or SC without uracil to saturation phase and serially diluted in sterile distilled H₂O to 10⁻⁴ and aliquots were streaked on solid media. The cells were cultured on YP, and YP supplemented with Na₃AsO₃ (As(III) (1 mM)), CdCl₂ (Cd (0.1 mM)), NiCl₂ (Ni (8 mM)), as well as 6% ethanol and 6% ethanol + Na₃AsO₃ (As(III) (0.3 mM)) for two days at 30°C. YP + 2% glucose (YPD) was used for the experiments without the involvement of any plasmid and YP + 2% galactose (YPG) was used for experiments harboring plasmids (overexpression) with a galactose-inducible promoter (GAL1/10), in order to activate the desired gene. The number of colonies under drug conditions were compared to the number of colonies formed under non-drug conditions for consistency and subsequently normalized to the number of colonies formed by wild type (WT) strain under the same condition. Each experiment was repeated at least three times.

**Quantitative β-galactosidase assay**

*ortho*-Nitrophenyl-β-galactoside (ONPG)-based β-galactosidase analysis was used to quantify involvement of the identified genes in the IRES-mediated translation of URE2 via quantification of β-galactosidase activity produced by a plasmid containing URE2-IRES fused to a β-galactosidase reporter (p281-4-URE2) [27]. This plasmid contains a DNA sequence that forms four strong hairpin loops prior to URE2-IRES region at its mRNA level. These four hairpin loops inhibit cap-dependent translation of URE2 mRNA. A background plasmid (p281) carrying only a β-galactosidase reporter was used as a control for cap-dependent translation [27]. Both plasmids contained a GAL1/10 promoter and YPG was used as a medium to activate the desired gene [32]. Each experiment was repeated at least three times.
Reverse Transcriptase quantitative PCR (RT-qPCR)

This methodology was used to assess the content of target mRNAs. Total mRNA was reverse-transcribed into complementary DNA (cDNA), using iScript Select cDNA Synthesis Kit (Bio-Rad) according to the manufacturer’s instructions. cDNA was then used as a template for quantitative PCR (qPCR). Total RNA extractions were performed with Qiagen RNeasy Mini Kit (Qiagen). qPCR was carried out using Bio-Rad iQ SYBR Green Supermix and the CFX connect real time system (Bio-Rad), according to the manufacturer’s instructions. In this experiment, PGK1 was used as a constitutive housekeeping gene and related to WT [31], [32]. Each qPCR experiment was repeated at least three times using separate cDNA samples. Following oligonucleotide primers were used to quantify URE2 and PGK1 mRNAs:

Forward URE2: ATGATGAATAACAACGGCAA
Revers URE2: TCATTCACCACGCAATGCC T
Forward PGK1: ATGTCTTTATCTTCAAAGT T
Revers PGK1: TTATTTCTTTTCGGATAAGA

Immunoblotting

Western blotting was used to quantify relative protein levels. Total protein was isolated using detergent-free methods as described in [33]. Samples were grown overnight in liquid YPD, pelleted and washed with PBS buffer. Samples for As(III) treatment, were grown overnight in liquid YPD and then treated with Na3AsO3 (As(III) (0.5 mM)) for 2 hours. Bicinchoninic acid assay (BCA) (Thermofisher) was used to quantify total protein concentration according to manufacturer’s instructions. Equal amounts of total protein extract (50 μg) were loaded onto a 10% SDS-PAGE gel, run on Mini-PROTEAN Tetra cell electrophoresis apparatus system (Bio-Rad) [34]. Proteins were transferred to a nitrocellulose 0.45 μm paper (Bio-Rad) via a Trans-Blot Semi-Dry Transfer (Bio-Rad). Mouse monoclonal anti-GFP antibody (Santa Cruz) was used to detect protein level of Ure2p in Ure2-GFP protein fusion strains. Pgk1p was also used as a constitutive housekeeping protein for quantification purposes. Mouse anti-Pgk1 (Abcam) was used to detect Pgk1p levels [31]. Immunoblots were visualized with chemiluminescent substrates (Bio-Rad) on a Vilber Lourmat gel doc Fusion FX5-XT (Vilber). Densitometry analysis was carried out using the FUSION FX software (Vilber), and each density was normalized to the density formed by Pgk1p control. Each experiment was repeated at least three times using three separate total protein isolates.

Polyribosome fractionation

The total RNA extraction from yeast cells was adapted from [35]. Overnight cell culture was used to inoculate YPD liquid medium. Prior to extraction, cycloheximide (100 μg/ml) was added to the samples for 15 minutes. Yeast culture was harvested at mid-log phase (OD600 0.6–0.8) and immediately chilled on dry ice prior to lysing the cells. Yeast cells were lysed by mechanical disruption using 425–600 μm acid-washed glass beads (Sigma). Fifty μg of the total RNA was then loaded on a 10–50% sucrose gradient (20 mM Tris pH 8, 140 mM KCl, 5 mM MgCl2, 0.5 mM DTT, 100 μg/ml cycloheximide and sucrose to desired concentration). An Automated Gradient Maker (Biocomp gradient maker) was used to produce the sucrose gradients. Centrifugation was performed at 40,000 rpm for 2 hours at 4°C (Beckman Optima LE-80K Ultracentrifuge) to separate the particles according to relative sedimentation rate.
Samples were analysed via a Biocomp Gradient Station immediately after centrifugation. The instrument recorded \( A_{254} \) using a flow cell coupled with a spectrophotometer (Bio-Rad Econo UV monitor). In this procedure, untranslated mRNAs (top fractions) are separated from polysome-associated mRNAs (bottom fractions) as described in [36]. Fractions were collected (~650 μl) using Bio-Rad Collection Station (Bio-Rad®) and adjusted to 1% SDS for fluctuation analysis of \( URE2 \) mRNA level via RT-qPCR [37]. Each polysome profiling experiment was repeated at least three times.

Luciferase RNA (0.1 μg/ml) (Promega®) was then added to each fraction as a control. RNA was precipitated overnight and purified as described in [37], by using Glycoblue and acidic Phenol/chloroform (pH 4). Purified RNA samples were subjected to quantification for \( URE2 \) mRNA by performing RT-qPCR as described in the above section. The normalized values for each fraction was determined by using the Ct values for \( URE2 \) and the Ct values for the gene of interest using the formula \( 2^{(Ct_{\text{luciferase}} - Ct_{\text{target gene}})} \) [38]. The relative amount of \( URE2 \) mRNA was calculated by dividing the amount in each fraction by the total signal in all fractions [38], [39]. RT-qPCR analysis for polysome profiling was repeated at least three times using fractions from three separate polysome profiles.

**Genetic interaction (GI) and conditional GI analysis**

Synthetic genetic array (SGA) analysis was performed in large-scale through the creation of double-mutations and subsequent analysis of colony size (fitness) as previously described in [31], [40]. In summary, both gene candidates \( ITT1 \) and \( RPS1A \) were knocked out in MATα (BY7092) and crossed with two arrays of haploid MATα knock-out strains [31]. The first array contained 384 deletion strains for genes that are directly or indirectly involved in the process of translation (translation gene deletion array). The second contained 384 random genes that were selected from the YKO collection [23], [31], which was used as a control. Selectable markers designated in each background mating type, allowed for multiple selection steps. Meiotic progeny harboring both mutations were selected. The created arrays could then be used to score double mutants for their altered fitness under certain conditions [23].

Colony size of both single mutant arrays (reference and control) and double mutant arrays were measured for their colony fitness [41], [42]. After three repeats, the interactions with 20% alteration or more in at least two repeats were considered positive hits. Conditional SGA analysis was carried on under sub-inhibitory concentrations of chemicals (0.7 mM for As(III) and 60 ng/ml for cycloheximide). Phenotypic suppression array (PSA) analysis infers closer functional relationships between interacting genes by establishing a more direct form of GI. It refers to situations where defective growth caused by deletion of a particular gene, in a specific growth condition (for example presence of a drug) is rescued by the overexpression of another. PSA analysis was performed under a high sub-inhibitory targeted condition (As(III) (1.2 mM) and cycloheximide (100 ng/ml)) as previously described in [31], [43], by overexpressing either \( ITT1 \) or \( RPS1A \) in the above described translation gene deletion array. Each experiment was repeated three times and the interactions with 25% alteration or more in at least two screens were scored as positive.

**Results and discussion**

**Deletion of \( ITT1 \) or \( RPS1A \) increases yeast sensitivity to heavy metals**

Understanding the biology of the stress that heavy metals exert on a cell, as well as the cellular responses and mechanisms that a cell uses for detoxification of these toxins, has been the subject of numerous investigations over the past decades e.g. [2], [14], [15], [16]. Although much has been learned, additional studies are needed to uncover details of such responses as well as
additional genes that may participate in this process. To this end, while screening for yeast gene deletion mutants against heavy metals, we identified two deletion mutant strains for ITT1 and RPS1A that showed increased sensitivity to three heavy metals (As, Ni and Cd). These two genes are part of a larger investigation to study the chemical-genetic profile of heavy metals in yeast e.g. [44]; details of this study will be published elsewhere. In our spot test sensitivity analysis, when As(III) (1 mM), Cd (0.1 mM) and Ni (8 mM) were added to the solid media the number of normalized yeast colony counts were significantly reduced for itt1Δ and rps1aΔ strains (Fig 1A), highlighting the sensitivity of the mutant strains to these metals and metalloids. Rescue experiments revealed a complete recovery to heavy metal toxicity upon the reintroduction of the deleted genes into their corresponding mutants (Fig 1B). This reversion of sensitivity indicates that the sensitivity phenotypes are in fact a consequence of the intended gene deletions and are not due to a possible secondary mutation within the genome. The sensitivity of deletion mutants to As, Cd and Ni suggests a potential association for the target genes to heavy metal toxicity, a unique observation that has not been previously reported.

Since the function of both ITT1 and RPS1A can be linked to the process of protein biosynthesis [21], [22], it is conceivable that these genes may indirectly influence heavy metal sensitivity by regulating the activity of another gene. However, given that the Ure2p is reported to be a key enzyme involved in heavy metal detoxification in yeast [19], [20], we made double gene deletion mutants for ITT1 and RPS1A with URE2 and exposed them to As(III) (1 mM) for further analysis (Fig 1C). The sensitivity analysis of the double gene deletion mutants indicated no increased sensitivity to As(III) in addition to that observed for the single gene deletion mutant for URE2. This specifies a dominant effect for URE2 on heavy metal sensitivity over ITT1 and RPS1A (Fig 1C). To further support the observed phenotype, we introduced plasmid vectors containing ITT1 and RPS1A into the deletion strain of URE2 (Fig 1D). The results demonstrated the same levels of sensitivity to As(III) with no compensation, deeming indirect roles of ITT1 and RPS1A in rescuing the cells from As(III) toxicity when URE2 is deleted. One way to explain this data is that ITT1 and RPS1A may exert their effect on sensitivity via the same pathway as URE2. If ITT1 and RPS1A influenced a second pathway, it might be expected that their deletion would have had an additional effect on sensitivity when combined with URE2 deletion [44]. However, this was not observed. On the other hand, overexpression of URE2 in itt1Δ and rps1aΔ strains reversed the As(III) sensitivity phenotype observed by the corresponding gene deletions, effectively deeming these gene deletions inconsequential for heavy metal sensitivity (Fig 1E). These observations are in accordance with the activity of URE2 as a dominant player in heavy metal toxicity and that it functions downstream of ITT1 and RPS1A. Finally, to ensure the specific sensitivity of itt1Δ and rps1aΔ strains to heavy metals, we investigated the introduction of different oxidative stress conditions such as methyl methanesulfonate (MMS), acetic acid, and heat shock (S1 Fig).

Our findings suggest that the influence of ITT1 and RPS1A on heavy metal sensitivity is linked to URE2. Although both ITT1 and RPS1A have reported roles in protein biosynthesis [21], [22], the possible mechanism, regulation of transcription or translation of URE2 remains to be investigated.

### ITT1 and RPS1A do not affect the expression of URE2 at the mRNA level

RT-qPCR was employed to detect possible changes to URE2 mRNA levels in the absence of ITT1 and RPS1A. Fig 2 illustrates our observation that in comparison to WT, the URE2 mRNA appeared unchanged in the mutant strains, itt1Δ and rps1aΔ. We also investigated the content of URE2 mRNA after exposure to As(III) (0.5 mM). As noted before, neither the deletion of
ITT1 or RPS1A appeared to influence the content of URE2 mRNA, suggesting that the activity of ITT1 and RPS1A is independent of URE2 mRNA content.

Ure2p content is reduced in the absence of ITT1 or RPS1A

We investigated the level of Ure2p in the presence and absence of ITT1 or RPS1A in the cells by Western blot analysis. This was accomplished using a strain that carried Ure2p fused to a GFP at the genomic level. Our analyses show a reduction in the endogenously-expressed Ure2-GFP fusion protein levels in the absence of ITT1 or RPS1A (Fig 3). When either ITT1 or RPS1A were deleted, Ure2p levels were reduced by approximately 40% and 60%, respectively, compared to the WT cells (Fig 3A). This suggests that ITT1 and RPS1A play an imperative role in regulating the expression of Ure2p. In parallel, introduction of As(III) (0.5 mM) to the growth media reduced Ure2p approximately 70% and 80% for Δitt1 and Δrps1a, respectively compared to WT strain (Fig 3B) which is reduced by approximately 34% in the presence of As (III) (0.5 mM) compared to normal experimental condition. Deletion of ITT1 or RPS1A did not change the protein levels of Pgp1p, used as an internal control. These observations provide evidence by connecting ITT1 and RPS1A activities to the level of Ure2p. The additional
reduction in the Ure2p levels in the presence of As(III) implicates that ITT1 and RPS1A may have a higher influence in regulation of Ure2p expression under a stress condition.

Analysis of the URE2 translation
Since our data suggests a role for ITT1 and RPS1A in modulating URE2 expression at the protein level, polyribosome-bound mRNA analysis was performed. In this method, fractions of polysomes are isolated and analyzed for their content of a target mRNA. Those mRNAs that are translated more efficiently are generally found in association with multiple ribosomes and hence will be isolated in heavier polysome fractions. In contrast, those mRNAs, which are translated to a lesser degree, can be found in lower density fractions [34]. In this way, the distribution of mRNAs within polysome fractions can be used to estimate the translation efficiency of the target mRNA. Using this strategy, polysome profile analysis was performed for URE2 mRNA, in the presence or absence of ITT1 and RPS1A. Analysis of the polysome fractions for URE2 mRNA content using RT-qPCR, normalized to a control (housekeeping) mRNA (PGK1), showed a shift in URE2 mRNA accumulation towards lighter polysome fractions for mutant strains (Fig 4), suggesting that when ITT1 or RPS1A are deleted, URE2 mRNA is translated to a less efficiently. These data provide direct evidence that ITT1 and RPS1A affect the translation of URE2 mRNA.
Ethanol increases As(III) sensitivity for ITT1 and RPS1A gene deletion strains

In addition to cap-dependent translation, URE2 mRNA has been shown to undergo a cap-independent translation, which represents an interesting mode of gene expression control [27]. Cap-dependent translation is mediated through the scanning of mRNA 5’ UTR to find a suitable start codon. In cap-independent translation, mRNA structures called Internal Ribosome Entry Site (IRES) mediate the interaction between ribosomes and the mRNA, independently of the 5’cap [45]. IRES-mediated translation is mainly used by RNA viruses but it can also be found in cellular mRNAs [46], [47]. The majority of translation in eukaryotes occurs through cap-dependent translation, whereas IRES-mediated translation is often associated with physiological conditions such as stress, where general translation is compromised [45], [48], [49].

Knowing that the exposure to heavy metal causes a stress condition for yeast cells e.g. [19], [50] we examined the possibility that ITT1 and RPS1A may influence IRES-mediated URE2 mRNA translation. As a quick measure to examine this possibility, ethanol sensitivity analysis was used as a positive stress control. Eukaryotic initiation factor 2A (eIF2A) acts as a regulator of IRES-mediated translation in S. cerevisiae cells [51]. Abundance of eIF2A is shown to specifically repress IRES-mediated translation of URE2 as well as other yeast genes with IRES forming region [51]. It was previously reported that ethanol reduces the expression of eIF2A at protein levels and hence promotes translation via IRES elements [51].

As described, URE2 plays a critical role in heavy metal detoxification [19], [49]. If ITT1 and RPS1A affect IRES-mediated translation of URE2 mRNA, one may expect the presence of ethanol to further enhance the sensitivity of itt1Δ and rps1aΔ to As(III) treatment. In this case,
presence of ethanol and deletion of either ITT1 or RPS1A could be considered to introduce a double effect on the same overall process.

As demonstrated in Fig 5, ure2Δ, itt1Δ and rps1aΔ showed hypersensitivity to a very low concentration of As(III) (0.3 mM) in the presence of 6% ethanol which suggests that ethanol increases the sensitivity of mutant strains to As(III). The mutant strains did not exhibit sensitivity to neither 0.3 mM of As(III) nor 6% ethanol separately. These data may suggest a connection for the activities of ITT1 and RPS1A to IRES-mediated translation. These outcomes are open to other explanations, for example, ethanol and As(III) may have alternative effects on the cell independent of IRES-mediated translation and hence influence cell viability. In this context double deletion strains for eIF2A with ITT1 or RPS1A showed similar sensitivity to the presence of either As(III) (0.3 mM) alone or with ethanol (6%) (Fig 5) connecting the reported activity of ethanol to eIF2A. Nonetheless, additional experiments are needed to confirm these interpretations. A double gene deletion for ITT1 or RPS1A with URE2 did not result in additional sensitivity compared to that observed for URE2 single gene deletion. This additionally reiterates the connection between the activities of ITT1 and RPS1A via URE2 expression.

**ITT1 and RPS1A affect IRES-mediated translation of a β-galactosidase reporter gene**

To further study the effect of ITT1 and RPS1A on IRES-mediated translation of URE2, we used a β-galactosidase reporter whereby the β-gal mRNA is under the translational control of the
URE2-IRES element. For this purpose, we utilized a previously published plasmid construct known as p281-4-URE2 (Fig 6) [27], [51].

As shown in Fig 6, deletion of either ITT1 or RPS1A resulted in reduced expression of β-galactosidase under the translational control of URE2-IRES. This data further supports the notion that ITT1 and RPS1A regulate URE2-IRES-mediated translation. As a control to account for cap-dependent translation activity, p281 background construct [27], [51] carrying a β-galactosidase mRNA lacking four hairpin loops and URE2-IRES was utilized. We observed no significant difference in β-galactosidase activity for WT, Δitt1 and Δrps1a strains indicating that ITT1 and RPS1A do not influence cap-dependent translation (Fig 6). Since the reduced levels of β-galactosidase may be a consequence of reduced mRNA, the β-galactosidase mRNA was evaluated using RT-qPCR. As expected, we observed no significant change in its mRNA level in the absence of neither ITT1 nor RPS1A (Fig 6B).

Genetic interaction analysis further connects the activity of ITT1 and RPS1A to regulation of translation in response to stress

To further examine the role of ITT1 and RPS1A in the process of translation, we studied the genetic interactions (GIs) by screening ITT1 and RPS1A against an array of 384 genes associated with protein biosynthesis and a second set of 384 random genes used for control purposes. Genes that are functionally related often partake in GIs (also known as epistatic interactions) [52], [53]. The most commonly studied form of GI is known as negative GI, where the reduced fitness or lethal phenotype of a double mutant strain for missing two genes is not observed in single mutant strains [31]. Genes that are associated with parallel and compensating pathways are thought to commonly form negative GIs [31], [40]. Under standard laboratory growth conditions, both ITT1 and RPS1A exhibited negative GIs with a limited number of genes involved in the process of protein biosynthesis (Fig 7A and 7B). This is expected as both genes are thought to be involved in the process of translation. Examples of
genes that formed GIs with ITT1 and RPS1A are large ribosomal subunit protein 19A (RPL19A) and general control non-derepressible (GCN3), respectively. Rpl19p is a conserved large ribosomal subunit protein involved in ribosomal intersubunit bridging and its alteration is connected to the fidelity of translation [54], [55]. On the other hand, GCN3 is the alpha subunit of translation initiation factor 2B (eIF2B), which is involved in guanine-nucleotide exchange for eIF2. In this manner, phosphorylation of eIF2B regulates the activity of Gcn3p [56], [57].

Conditional GIs define an interesting type of GIs as they provide more compelling insight on the function of target genes under specific conditions [31], [58]. They represent the mosaic nature of gene function(s) which can change as a result of different external or internal factors. For example, although a number of genes are known to play a role in the DNA repair pathway, their expression is only regulated in response to the presence of DNA damage [29], [43]. We therefore investigated the negative GIs by assaying ITT1 and RPS1A in the presence of a low concentration of As(III) (0.7 mM). Under this condition, we found both genes to form a new set of interactions with a series of genes that play a role in the regulation of translation (Fig 7A and 7B). This data suggests that in the presence of As(III), both genes appear to gain a new role in regulating the process of translation (p-value ITT1 = 9.39e-5 and p-value RPS1A = 5.26e-10) (Fig 7A and 7B). Suppressor of ToM1 (STM1) is an example of a gene that formed new negative genetic interactions in the presence of As(III) with both ITT1 and RPS1A. STM1 codes for a protein that is required for optimal translation under nutrient stress [59], [60]. Enhancer of mRNA DeCapping EDC1 and its paralog EDC2 are other examples of the negative interactions gained by RPS1A under As(III) condition. Edc proteins directly bind to mRNA substrates and activate mRNA decapping. They also play a role in translation during stress conditions such as heat shock [61]. We also studied negative GIs in the presence of cycloheximide, which binds to the E-site of the 60S ribosomal subunit and interferes with deacetylated
tRNA to inhibit general protein synthesis in the cell [62] (Fig 7A and 7B). Similar to our observations with As(III), in the presence of a mild concentration of cycloheximide (60 ng/ml), ITT1 and RPS1A formed new negative GIs with a group of genes that are associated with the regulation of translation (p-value \(\text{ITT1} = 1.33e-6\) and p-value \(\text{RPS1A} = 6.59e-9\)). Translation Initiation Factor 4A (TIF2) is an example of the gained interaction for both ITT1 and RPS1A in response to cycloheximide. TIF2 is a key player in translation initiation and holds a helicase activity [63]. Altogether, these observations are in agreement with the involvement of ITT1 and RPS1A in regulating translation in response to stress.

To further study ITT1 and RPS1A, we conducted phenotypic suppression array (PSA) analysis [31], [65] (Fig 7C and 7D) to assess the ability of overexpression of our target genes to reverse the defective phenotype (i.e. sensitivity) on a series of gene deletion strains under specific conditions. PSA analysis constitutes a more direct form of GI and can infer close functional relationships between interacting genes. In these cases, overexpression of one gene compensates the phenotypic adverse effect that is caused by the absence of another gene under certain conditions, such as stress caused by different chemicals [30], [31]. We observed that the overexpression of both ITT1 and RPS1A reversed the sensitivity of a number of gene deletion strains to As(III) (1.2 mM) or cycloheximide (100 ng/ml) (Fig 7C and 7D). The majority

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Fig 7. Genetic interaction (GI) analysis for ITT1 and RPS1A. (A) Negative GIs for ITT1 under standard growth conditions and the presence of As(III) or cycloheximide. In this case, deletion of a second gene along with ITT1 forms an unexpected growth reduction. (B) Negative GIs for RPS1A under standard growth conditions and the presence of As(III) or cycloheximide as in (A). (C) Phenotypic suppression array (PSA) analysis using the overexpression of ITT1. In this way, overexpression for ITT1 compensated for the sensitivity of gene deletions to As(III) (1.2 mM) or cycloheximide (100 ng/ml). (D) PSA analysis using the overexpression of RPS1A as in (C). Each experiment was repeated three times and the interactions with 20% alteration or more in at least two screens were scored as positive. P-values were obtained from GeneMANIA [64].
of the newly identified gene interactors are involved in translation regulation further connecting the activity of ITT1 and RPS1A to the regulation of translation. Interestingly, two of these genes, GIS2 and DOM34, have reported to have IRES trans-acting factor (ITAF) activity [31], [66], [67]. The fact that the overexpression of ITT1 and RPS1A can compensate for the absence of ITAFs, GIS2 and DOM34 provide further evidence connecting ITT1 and RPS1A to IRES-mediated translation.

GIg Suppressor (GIS2) is a well studied translational activator for numerous IRES containing mRNAs [66], [67]. Duplication Of Multilocus region 34 (DOM34) is a protein that facilitates inactive ribosomal subunit dissociation to aid in translation restart [68], and is reported to play a role in IRES-mediated translation in yeast [31]. Our PSA analysis, not only supports a role in IRES-mediated translation pathway for both ITT1 and RPS1A, but also suggests a possible systematic compensation between certain ITAFs which can be the subject of future studies. This also leads to the conclusion, that other interacting partners of ITT1 and RPS1A in this experiment may play a role in IRES-mediated translation. Further studies are required to investigate these hypotheses. In agreement with our findings here, Sammons et al., [67] reported a physical interaction between Rpsa1p and Gis2p connecting the activity of these two proteins.

Supporting information

S1 Fig. Normalized CFU counts for different yeast strains after exposure to acetic acid (180 mM), heat shock (37˚C), and MMS (0.05% v/v). CFU counts after 3 days exposure to the experimental conditions are normalized to control condition counts. Sensitivity of itt1Δ, rps1aΔ are compared to WT phenotype in the presence of (A) acetic acid (180mM), (B) heat shock (37˚C) and (C) MMS (0.05% v/v). For each experimental condition, a corresponding positive control strain is included. Each experiment was repeated at least three times. Error bars represent standard deviations. * Indicates statistically significant differences (p <0.005).

S2 Fig. Spot test sensitivity analysis for different yeast strains. Cells were grown to saturation, serially diluted, and spotted on solid media containing As(III) (1 mM), Cd (0.1 mM), Ni (8 mM) or no drug. Plates were incubated at 30˚C for 3 days.

S3 Fig. Spot test sensitivity analysis of selected GIs. Cells were grown to saturation, serially diluted, and spotted on solid media with or without As(III). (A) and (B) EDC1 and DOM34 show conditional GI with ITT1 and RPS1A in the presence of As(III) (0.7 mM) representing conditional negative genetic interactions. DAN1 is used as a negative control. (C) and (D) Overexpression of ITT1 and RPS1A compensated the sensitivity of gene deletion strains for GIS2 and DOM34, respectively, in the presence of As(III) (1.2 mM) confirming a phenotypic suppression GI for ITT1 with GIS2 and RPS1A with DOM34. DAN1 was used as a negative control.

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