Genetic and Metabolomic Dissection of the Ergothioneine and Selenoneine Biosynthetic Pathway in the Fission Yeast, *S. pombe*, and Construction of an Overproduction System

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Abstract

Ergothioneine is a small, sulfur-containing metabolite (229 Da) synthesized by various species of bacteria and fungi, which can accumulate to millimolar levels in tissues or cells (e.g. erythrocytes) of higher eukaryotes. It is commonly marketed as a dietary supplement due to its proposed protective and antioxidative functions. In this study we report the genes forming the two-step ergothioneine biosynthetic pathway in the fission yeast, *Schizosaccharomyces pombe*. We identified the first gene, *egt1* (SPBC1604.01), by sequence homology to previously published genes from *Neurospora crassa* and *Mycobacterium smegmatis*. We showed, using metabolomic analysis, that the *Δegt1* deletion mutant completely lacked ergothioneine and its precursors (trimethyl histidine/hercynine and hercynylcysteine sulf oxide). Since the second step of ergothioneine biosynthesis has not been characterized in eukaryotes, we examined four putative homologs (NFs/SPBC21D10.11c, SPAC11D3.10, SPCC777.03c, and SPBC660.12c) of the corresponding mycobacterial enzyme EgtE. Among deletion mutants of these genes, only one (ΔSPBC660.12c, designated *Aegt2*) showed a substantial decrease in ergothioneine, accompanied by accumulation of its immediate precursor, hercynylcysteine sulf oxide. Ergothioneine-deficient strains exhibited no phenotypic defects during vegetative growth or quiescence. To effectively study the role of ergothioneine, we constructed an *egt1* overexpression system by replacing its native promoter with the *nmt1* promoter, which is inducible in the absence of thiamine. Thus, we employed three versions of the *nmt1* promoter with increasing strength of expression and confirmed corresponding accumulations of ergothioneine. We quantified the intracellular concentration of ergothioneine in *S. pombe* (0.3, 157.4, 41.6, and up to 1606.3 μM in vegetative, nitrogen-starved, glucose-starved, and *egt1*-overexpressing cells, respectively) and described its gradual accumulation under long-term quiescence. Finally, we demonstrated that the ergothioneine pathway can also synthesize selenoneine, a selenium-containing derivative of ergothioneine, when the culture medium is supplemented with selenium. We further found that selenoneine biosynthesis involves a novel intermediate compound, hercynylselenocysteine.

Introduction

Ergothioneine (EGT) is a sulfur-containing, Nα,Nα,Nα-trimethyl-L-histidine-derived metabolite that is synthesized by various species of bacteria and fungi; recently extensively reviewed by Cheah and Halliwell [1]. Higher organisms obtain EGT in food and accumulate it in certain tissues up to millimolar levels [2,3] through a specific transporter, ETT/OCTN1 [4]. In mammals, large amounts of EGT are found in erythrocytes, bone marrow, liver, kidney, eye lens, and seminal fluid [2,5,6]. Nevertheless, EGT is neither a nutrient (it is virtually unmetabolized in humans) nor a vitamin (it is non-essential). EGT is commonly marketed as a dietary supplement or nutraceutical, due to its anti-oxidant properties *in vitro*, reported in numerous publications [3,7–10]. Direct scavenging of free radicals and chelation of transition metals are the most widely cited possible functions of EGT [10,11]. However, so far, no rigorous research has conclusively demonstrated any benefit of EGT *in vivo*. It is unclear whether EGT consumption contributes to human health, and if it does, what daily intake is optimal. It is thus of considerable interest for biologists and medical scientists to uncover the physiological mechanism of EGT at the molecular level. Recently, biosynthetic pathways for EGT have been characterized in *Mycobacterium smegmatis* [12] and *Neurospora crassa* [13], allowing the use of genetic methods.

The fission yeast, *Schizosaccharomyces pombe*, is a suitable model organism for the study of cell division and quiescence [14–16]. We have previously established a method of comprehensive metabolomic analysis in *S. pombe* using liquid chromatography-mass spectrometry (LC-MS) [17]. Among the several hundred observed
metabolites, we also identified EGT and described its accumulation under glucose starvation [18] and nitrogen starvation [19]. In addition, we reported abnormally high accumulations of EGT in the proteasome regulatory subunit mutant mts3–1, which suffers from severe oxidative stress caused by mitochondrial dysfunctions in G0 arrest [20], and in the Kruppel-like zinc-finger transcription factor deletion mutant Δgfl1, which exhibits cell wall defects in long-term quiescence, accompanied by up-regulation of mitochondrial transcripts [21]. Interestingly, EGT is not synthesized by the budding yeast, Saccharomyces cerevisiae [22], suggesting that not all fungi require EGT for normal cellular function. Due to its simplicity and advanced genetics, S. pombe might represent an ideal unicellular, eukaryotic system to study the biochemical role of EGT.

Here we report identification of genes forming the two-step biosynthetic pathway of EGT from histidine in S. pombe, through the combined use of genetic and metabolomic approaches. Among other compounds, we were able to identify the direct precursor of EGT, hercynylcysteine sulfoxide. We constructed an EGT and hercynylcysteine sulfoxide overproduction system utilizing three different overexpression strains with the inducible mnt1 promoter. We quantified intracellular EGT content in wild type (WT) as well as overexpression cells by constructing a calibration curve from pure EGT standard injections into the LC-MS. Further, we show the accumulation of EGT under long-term quiescence. Finally, we demonstrate that the EGT pathway can also synthesize seleno-enzyme, a selenium-containing derivative of EGT, the production of which involves a novel intermediate compound, hercynylselenocysteine.

**Results**

**EGT biosynthesis pathway**

The reported EGT biosynthetic pathways in M. smegmatis and N. crassa are schematized in Figure 1A. On the basis of sequence homology, the S. pombe locus, mug158'/SPBC1604.01, was previously suggested to encode the main EGT biosynthetic enzyme [12,13]. This enzyme, with 773 amino acids, catalyzes triple methylation of histidine to hercynine (Nɛ,Nɛ,Nɛ-trimethyl histidine) and subsequent conjugation with cysteine and oxygen to form hercynylcysteine sulfoxide. mug158'/SPBC1604.01 is a distant homolog of the mycobacterial EgdD and EgdB genes, encoding a single fusion protein. Figure 1B shows the domain structure of this protein, according to the Conserved Domain Database [23], in comparison with its homologs in Schizosaccharomyces japonicus (a relative of S. pombe), N. crassa, and M. smegmatis. The exact locations and sequences of individual domains are shown in Table S1 and the amino acid sequence alignment is shown in Table S2. The exact locations and sequences of conserved domains in Table S2; amino acid sequence alignment in Figure S2). Among these, SPAC11D3.10, and SPCC777.03c have highly similar sequences, possibly originating from horizontal gene transfer (Figure 1D).

To identify the EgdE homolog in S. pombe, we obtained deletion mutants from the Bioneer haploid deletion library [24], cultivated them under nitrogen starvation (EMM2 medium lacking NH4Cl, hereafter designated EMM2-N), for 24 hours to induce EGT production, and performed a metabolomic analysis. Among the four deletion mutants tested, ΔSPBC660.12c was the only one showing a substantial decrease in EGT and an increase in its precursor, hercynylcysteine sulfoxide (Figure 1E; numerical results of all LC-MS measurements shown in Table S3). We thus designated the SPBC660.12c locus egt2⁺, as it represents the gene primarily responsible for the second step of EGT biosynthesis in S. pombe.

**Verification of egt1⁺ and egt2⁺ by metabolomic analysis**

To verify correct assignment of the egt1⁺ and egt2⁺ genes, we newly constructed full deletion mutants, Δegt1 and Δegt2, by replacing the target loci in the WT k⁺ 972 strain with the kanamycin resistance marker (kanMX). Correct integration of the kanMX modules into the new strains was verified by PCR. No difference from WT in cell size or shape was observed in these strains under any of the analyzed conditions. Table 1 shows the results of metabolome analysis of the constructed strains under starvation (EMM2-N or EMM2 medium with low concentration of glucose, 1.1 mM, hereafter designated EMM2-LG, for 24 hours). Data regarding EGT and its precursors, starting with histidine, are shown. We confirmed the absence of all pathway intermediates in Δegt1 and the accumulation of hercynylcysteine sulfoxide in Δegt2. A small amount of EGT was still found in Δegt2 under starvation. To determine whether any other cysteine desulfurase might contribute to this enzymatic reaction, we constructed multiple deletion mutants of SPBC660.12c (egt2⁺) with the other candidate EgdE homologs, but even in successfully constructed double and triple mutants, a significant amount of EGT still remained (Figure S3). This, however, is not a surprising result, as Scebeck [12] previously demonstrated that hercynylcysteine sulfoxide could spontaneously convert into EGT in the presence of PLP. Furthermore, as also shown by Scebeck [12], this reaction could be catalyzed by an unrelated PLP-binding β-lyase originating from Erwinia tasmaniensis. Thus, we suspect the residual EGT found in Δegt2 might be a product of a hercynylcysteine sulfoxide reaction with PLP, possibly catalyzed by an unrelated PLP-binding enzyme (S. pombe genome contains at least 26 PLP-binding enzymes, according to PomBase [25]).

Neither the Δegt1 nor the Δegt2 strain showed any growth defects during cultivation in either rich (YE) or minimal (EMM2) culture media. Furthermore, deletion of egt1⁺ or egt2⁺ caused no significant perturbation to the intracellular metabolome of quiescent cells (Figures 2A and 2B). Apart from the disappearance of EGT and its precursors, all other metabolite levels remained within the range of common experimental error.

**Conservation of egt1⁺ and egt2⁺ in other species**

As observed by Scebeck [12], the EGT biosynthetic pathway can be found in a relatively small number of eukaryotes, notably in the phyla Basidiomycota (most species) and Ascomycota (most species in subphyla Pezizomycotina and Schizosacharomycetes). We summarized homologs of EGT biosynthesis genes in several organisms closely related to S. pombe (Table 2). The exact locations of the conserved domains in these enzymes are shown in Tables S4 and S5. According to sequence alignment (Figures S4 and S5),
conserved domains show higher homology among species than inter-domain regions. Methyltransferase and FGE-sulfatase domains contain long non-homologous sequence inserts in *U. maydis* UM00197. The function of the inserts in these two proteins is unknown, however, they are not unusual among various species [13].

Based on sequence homology to *S. pombe* Egt2 (33% amino acid sequence identity), we propose that the NCU11365 gene may encode the second enzyme for EGT biosynthesis in *N. crassa*. Bello et al. [13] previously suggested the NCU04636 gene (19% identity to Egt2), which appears to correspond to nfs1, the mitochondrial cysteine desulfurase in *S. pombe* (63% identity). Both *S. pombe* nfs1 and *N. crassa* NCU04636 appear to be the closest homologs of mycobacterial EgtE (as also indicated in Figure 1D). However, we did not detect any change in hercynylecystine sulfoxide levels in the Δnfs1 deletion mutant (Figure 1E). Furthermore, the nfs1 gene has apparent homologs in species that do not produce EGT, such as *S. cerevisiae* or humans (both genes called NFS1).

**Overexpression of egt1**

To study the overproduction effect of EGT in *S. pombe*, we applied the method of Bahler et al. [26] to replace the egt1 native promoter with the nmt1 promoter, which is inducible in the absence of thiamine [27]. We employed three versions of the nmt1 promoter plasmid with increasing strength of expression and constructed three strains P81nmt1-egt1+, P41nmt1-egt1+, and P3nmt1-egt1+, respectively. Using metabolomic analysis we confirmed the accumulation of EGT and its precursors in these strains, and this accumulation was effectively suppressed by the addition of 5 μg/ml thiamine to the EMM2 medium (Figure 2C).

**Intracellular EGT content**

Wild type *S. pombe* cells contain only trace amounts of EGT under normal vegetative conditions (Table 1). However, EGT increases in quiescent cells under starvation [18,19]. We measured the areas of EGT peaks under vegetative, quiescent, and egt1 overexpressing conditions, and converted them to absolute concentrations using a calibration curve based upon pure EGT injections ranging from 1 fmol - 10 nmol (Figure S6). Intracellular volume was assumed to be 148.5 μm3 for vegetative cells [28]. For nitrogen- and glucose-starved cells, intracellular volumes were estimated as 1/3 and 2/3 of the vegetative cell volume, respectively. The resulting intracellular concentrations are shown in Table 3.

To assess long-term variation in EGT content, we measured the level of intracellular EGT using metabolomic analysis during a 20-day time course under quiescence induced by nitrogen (EMM2-N) and glucose (EMM2-LG) starvation, respectively (Figure 2D). Cells showed time-dependent accumulation of EGT, despite being deprived of nutrients, suggesting that EGT might support cellular health under long-term quiescence. However, no loss of viability was observed in Δegt1 mutant during 20 days of starvation (Figure 2E).

**Table 1.** Normalized peak areas of the four compounds composing the EGT biosynthetic pathway obtained by metabolomic analysis of WT and newly constructed strains.

| Compound, peak m/z and retention time/Strain, cultivation condition | Histidine | Trimethyl-histidine (hercynine) | Hercynyl-cysteine sulfoxide | Ergothioneine |
|------------------------|-----------|---------------------------------|-----------------------------|--------------|
| EMM2                  | 156.077 m/z @12.4 min | 198.124 m/z @10.3 min | 333.123 m/z @12.2 min | 230.096 m/z @12.6 min |
| WT 972                | 14.4      | 1.7                            | 0                           | 0.1          |
| EMM2-N (24 h)         | 2.2       | 3.2                            | 2.2                         | 13.7         |
| EMM2-LG               | 55.4      | 65.9                           | 3.5                         | 6.7          |
| EMM2                  | 10.7      | 0.3                            | 0                           | 0            |
| Δegt1                 | 2.8       | 0.1                            | 0                           | 0            |
| EMM2-N (24 h)         | 54.1      | 0                              | 0                           | 0            |
| EMM2                  | 11.4      | 0.9                            | 3.9                         | 0            |
| Δegt2                 | 1.9       | 1.8                            | 58.2                        | 3.1          |
| EMM2-LG               | 61.1      | 64.7                           | 44.1                        | 1.6          |

Values were measured from metabolome samples of four different *S. pombe* strains in three different cultivation conditions, as indicated. Mass values (m/z) and LC retention times (min) of each peak are included for reference.

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Figure 2. Characterization of Δegt1 and Δegt2 strains. A and B. A scatter plot comparing results of metabolomic analysis of WT vs. Δegt1 (A) or Δegt2 (B) strains under nitrogen starvation (24 h in EMM2-N medium). Each dot represents a single identified metabolite. Values on both scales indicate normalized peak areas of metabolite peaks in corresponding strains. Red diagonal lines indicate a 2-fold difference. C. Results of metabolomic analysis of WT and egt1+ overexpression strains. Cells were cultivated at 26°C in the EMM2 medium lacking thiamine for at least 24 h. Cultures indicated +Thiamine were cultivated in the presence of 5 μg/ml thiamine for 24 h. Normalized peak areas of compounds composing the EGT pathway are shown. D. Time course metabolomic analysis of quiescent S. pombe cultures under nitrogen (EMM2-N) and glucose (EMM2-LG) starvation. Values represent means ± standard deviations of normalized peak areas of EGT in three independent cell cultures. E. Time course viability results of WT and Δegt1 deletion mutant cultures under nitrogen (EMM2-N) and glucose (EMM2-LG) starvation.

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Table 2. Closest homologs of S. pombe Egt1 and Egt2 proteins in selected species.

| Organism                          | Closest homolog of S. pombe Egt1 | Closest homolog of S. pombe Egt2 |
|-----------------------------------|----------------------------------|---------------------------------|
| Schizosaccharomyces japonicus     | SJAG_00832                       | SJAG_03856                      |
| Schizosaccharomyces octosporus    | SOCG_01424                       | SOCG_02548                      |
| Neurospora crassa                 | NCUI04343 (NcEgt-1)              | NCUI1365                        |
| Aspergillus niger                 | An19g05880                       | An02g02030 or An05g02190        |
| Aspergillus oryzae                | Ao090012000265                   | Ao090026000291                  |
| Ustilago maydis                   | UM00197                          | UM04128                         |

Candidate homologs were searched using the on-line version of the Basic Local Alignment Search Tool (http://blast.ncbi.nlm.nih.gov) for protein sequences (blastp) and candidates with the best similarity scores (lowest blastp E-values) were selected.
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Contribution of egt1+ to oxidative stress response

As EGT is generally considered to be a physiological antioxidant, and the ΔNcEgt-1 mutant was reportedly sensitive to tert-butyl hydroperoxide in N. crassa [13], we performed spot test experiments on EMM2 plates containing oxidants hydrogen peroxide and tert-butyl hydroperoxide, using the deletion and overexpression mutants described above (Figure S7). However, we did not observe any sensitivity or resistance of these strains compared to WT 972 strain, suggesting that egt1+ might not be among the primary mechanisms that protect S. pombe from exogenous peroxide.

 Biosynthesis of selenoneine

Selenoneine (Figure 3A) is a selenium-containing derivative of EGT found in tuna, and possibly implicated in methylmercury detoxification [29,30]. As the EMM2 medium does not normally contain selenium, it is not surprising that no selenoneine was detected in previous S. pombe metabolome data sets. To test whether S. pombe can produce selenoneine, we first examined cell cultivation in a liquid EMM2 medium supplemented with various concentrations of Na2SeO4 (Figure S8). The medium with 10 μM Na2SeO4 [hereafter designated EMM2+Se] was selected for further experiments, as S. pombe cells exhibited quite normal (albeit slightly slower) proliferation in this condition. Metabolomic analysis was performed on WT vegetative cells, WT nitrogen-starved cells (24 h in EMM2-N+Se), and P3nmt1-egt1+ overexpression mutant cells. We could clearly observe accumulation of selenoneine in the P3nmt1-egt1+ strain cultivated in EMM2+Se medium (Figure 3B), suggesting that the overexpressed egt1+ gene was also responsible for selenoneine synthesis. A tiny signal of selenoneine (<1% of the EGT signal intensity) could also be detected in WT nitrogen-starved cells, and the signals of both EGT and selenoneine disappeared in the Δegt1 deletion mutant (Figure 3C).

To rule out the possibility that selenoneine was produced by direct conversion from ergothioneine (without requiring any egt1+ activity), we performed two additional experiments. First, pure EGT was mixed with an equal concentration of Na2SeO4 in vitro and incubated at room temperature for 24 h. No selenoneine signal was detected in this mixture (Figure S9). In the second experiment we supplemented the Δegt1 mutant with 1 mM EGT, which was apparently transported into the cells and produced a strong EGT peak. No selenoneine was detected in this case either (Figure 3D). Furthermore, the weak, but clearly detectable selenoneine signal found in WT cells did not increase as a result of EGT supplementation. These results suggest that egt1+ activity is indispensable for selenoneine biosynthesis in S. pombe.

As we did not observe any signal of the presumed intermediate, hercynyl-selenocysteine sulfoxide, we constructed a double mutant of P3nmt1-egt1+ and Δegt2. This mutant should accumulate large amounts of the intermediate, assuming that the Egt1/Egt2 pathway is used for selenoneine synthesis. Surprisingly, we found a strong signal of hercynylselenocysteine (not sulfoxide) in this mutant (Figure 3E). This signal was also found in the P3nmt1-egt1+ single mutant, but further increased in the double mutant with Δegt2. We thus conclude that selenoneine biosynthesis, unlike EGT biosynthesis, does not produce a sulfoxide as its intermediate, but produces hercynylselenocysteine instead.

Since selenium naturally occurs in a very characteristic set of isotopes, we verified the identities of selenoneine and hercynylselenocysteine peaks by checking their isotope distribution patterns (Figure S10). Finally, to check whether the EGT/selenoneine pathway could possibly be involved in selenium detoxification in S. pombe, we performed a spot test experiment on EMM2 plates

Table 3. Absolute intracellular EGT concentrations (µM) in S. pombe cells.

| Cell condition   | Culture medium | Intracellular EGT (µM) |
|------------------|----------------|------------------------|
| WT vegetative    | EMM2           | 0.3                    |
| WT nitrogen starvation | EMM2-N (24 h) | 157.4                  |
| WT glucose starvation | EMM2-LG (24 h) | 41.6                   |
| P81nmt1-egt1+    | EMM2           | 32.4                   |
| P41nmt1-egt1+    | EMM2           | 181.2                  |
| P3nmt1-egt1+     | EMM2           | 1606.3                 |

Intracellular concentrations were derived from measured normalized peak areas using a calibration curve generated by injections of pure EGT in 10-fold dilution steps.

The detailed calculation method is described in Figure S6.
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Figure 3. Production of selenoneine in *S. pombe*. A. Chemical structure of selenoneine. B. Results of metabolomic analysis of WT and P3nmt1-egt1 strains in EMM2+Se and EMM2-N+Se media. Normalized peak areas of EGT and selenoneine are shown. C. Extracted ion chromatograms of EGT and selenoneine masses in raw LC-MS data acquired from metabolome samples of nitrogen-starved cells (24 h in EMM2-N+Se medium) of WT and Δegt1 strains. Note that the intensity scale of the selenoneine plot is 1% relative to that of the EGT plot. D. Extracted ion chromatograms of EGT and Ergothioneine in Fission Yeast PLOS ONE | www.plosone.org 7 May 2014 | Volume 9 | Issue 5 | e97774
containing various concentrations of Na₂SeO₄. However, none of the analyzed mutants showed any growth differences compared with WT cells (Figure S11).

**Discussion**

In this study we applied metabolomic analysis to identify the egt1⁺ and egt2⁺ genes composing the EGT and selenoneine biosynthetic pathway in *S. pombe* (Figure 4). These two genes have no homologs in *S. cerevisiae*, consistent with the fact that budding yeast do not produce EGT. In the future, a comparison between these two yeasts that appear similar, but are genetically rather distant, might provide useful clues regarding the native physiology of EGT or selenoneine. The presence or absence of the EGT pathway could be related to differences in the ecology of these two species.

In an environmental stress study, transcription of the egt1⁺ gene was up-regulated ~10-fold in the presence of 0.5 mM H₂O₂, and ~4-fold in the presence of Cd²⁺ [31]. It thus appears that egt1⁺ is strongly stress-responsive, providing support for the hypothesized antioxidant role of EGT. In addition, the egt1⁺ locus was previously named *mug158⁺*, denoting Meiotically Up-regulated Gene 158, due to its up-regulation upon entry into meiosis [32]. The egt1⁺ promoter contains the TR-box motif 5'-TTCTTTGTTY-3', recognized by the sexual development transcription factor, Ste11 [33], and a 5'-GTAAAYA-3'-binding motif recognized by the forkhead transcription factor, Mei4 [34]. Indeed, the egt1⁺ transcript was up-regulated in strains overexpressing ste11⁺ or mei4⁺ [35,36]. Also, egt1⁺ was up-regulated in a strain overexpressing both basic leucine zipper (bZIP) transcription factors *afl1⁺* and *afl31⁺* [36]. Furthermore, egt1⁺ was previously reported to be a putative regulatory target of the transcription factors, Ams2, Php5, and Pcr1, by analysis of multiple genome-wide microarray datasets [37]. Ams2 is a cell cycle-dependent GATA factor activated during replication [38]. Php5 is a subunit of the CCAAT-binding complex, which regulates respiration, based on glucose and iron availability [39]. Pcr1 is a CREB/ATF protein involved in stress response and sexual development [40,41]. In addition, the 5’ UTR region of egt1⁺ contains the CuSE element sequence, 5’-DWDDHGCTGD-3’, which is recognized by the Cuf1 transcription factor and activated by copper deficiency [42]. However, no transcriptional activation of egt1⁺ was found under varying copper levels [43], suggesting this 5’ UTR DNA fragment might not correspond to an actual Cuf1 binding site. In conclusion, it seems that egt1⁺ might be under regulatory control of a variety of different transcription factors.

In our study, the Δegt1 deletion mutant showed a complete absence of EGT and all its precursors. In the Δegt2 deletion mutant, some amount of EGT remained, consistent with results previously reported by Seebeck [12], that hercynylcysteine sulfoxide can spontaneously convert to EGT in the presence of PLP. Judging from the wealth of published transcriptome data, transcription of egt2⁺ does not vary in response to environmental conditions [31], and is only mildly up-regulated (~2-fold) during meiosis [32], suggesting that egt1⁺ represents the main regulatory

Figure 4. Summary of the described EGT and selenoneine biosynthetic pathway in *S. pombe*, and its transcriptional regulation.

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element in this pathway. The Egt2 enzyme could be ubiquitously present in the cells, simply converting any available hercynylcysteine sulfoxide into EGT.

Overexpression strains, P81nmt1-egt1, P41nmt1-egt1, and P5nmt1-egt1, showed gradual accumulation of EGT in standard EMM2 medium without thiamine, providing an additional confirmation of the correct egt1 assignment. Interestingly, P5nmt1-egt1 cells contained a large amount of hercynylcysteine sulfoxide. A possible explanation is that the activity of Egt1 in this strain was higher than the native activity of Egt2, resulting in accumulation of the metabolic intermediate between these two enzymes. These overexpression strains will be invaluable for verification of any proposed EGT mechanism. We also demonstrated that the P5nmt1-egt1 strain could synthesize a considerable amount of selenoneine when selenium was supplemented in the culture medium. Interestingly, selenoneine biosynthesis, unlike EGT biosynthesis, did not involve a formation of a sulfoxide intermediate, but rather involved a simple conjugate compound, hercynylselenocysteine. It is unknown whether egt1-mediated synthesis of hercynylselenocysteine requires oxygen, as in the case of EGT synthesis. The P5nmt1-egt1 strain proved to be very useful to elucidate the complete pathway, and we propose that this strain could also be employed industrially to produce EGT, as well as hercynylcysteine sulfoxide, hercynylselenocysteine, or selenoneine (the latter three compounds are not commercially available at present). The signal intensity of selenoneine found in WT nitrogen-starved cells was rather low; however, selenoneine might potentially have an interesting function in S. pombe. Selenoneine can be found in humans [44], and it was suggested that its radical-scavenging activity is even higher than that of EGT [30]. Importantly, we showed that selenoneine does not seem to be involved in detoxification of selenium in S. pombe.

From a wider perspective, the ubiquitous presence of EGT in living organisms, from bacteria to humans, suggests a crucially important function, yet this function has not been convincingly demonstrated so far. We found only trace amounts (0.3 μM) of EGT in vegetatively growing S. pombe cells; however, under starvation-induced quiescence it increased several hundred-fold. That, together with reported up-regulation of the egt1 transcript in the presence of H2O2 and in meiosis, implies a supportive or redundant systems to deal with oxidative stress, it is possible that the lack of EGT could be compensated by another mechanism. On the other hand, the observation that the highly overexpressed P5nmt1-egt1 strain, which accumulated EGT well beyond normal physiological levels, did not acquire any resistance to the tested oxidants, is intriguing. An attractive possibility exists that the true physiological purpose of EGT might lie in a yet unexplored area. We propose that genetic and metabolomic analyses, together with the collection of S. pombe strains introduced in this manuscript, may provide ideal tools to further investigate the in vivo role of this enigmatic compound.

Materials and Methods

Amino acid sequence alignment and bioinformatic analysis

Amino acid sequence alignment was performed using the online version (http://www.ncbi.nlm.nih.gov/tools/cobalt/) of the Constraint-based Multiple Alignment Tool (COBALT) [45] with default parameters. A phylogenetic tree was generated from the on-line COBALT tool (using the Fast Minimum Evolution method with Max Seq Difference set to 0.9 and other parameters set to default values) and visualized using Analyses of Phylogenetics and Evolution (APE) software [46]. Percentage identity of the amino acid sequences was calculated using the Clustal-Omega algorithm [47] with default parameters.

Strains and growth conditions

The S. pombe strains used in this manuscript are listed in Table 4. The synthetic minimal medium (EMM2), rich yeast extract medium (YE) and sporulation-inducing medium (MEA) recipes were used as published previously [48]. The following variants of the liquid EMM2 medium were used: EMM2-N (EMM2 lacking NH4Cl), EMM2-LG (EMM2 containing 1.1 mM – or 0.2 g/l – glucose), EMM2+Se (EMM2 containing additional 10 μM Na2SeO3), and EMM2+N+Se (EMM2-N containing additional 10 μM Na2SeO3). Cell cultures were cultivated at 26°C.

Construction of mutants

All DNA recombinant strains were constructed using a two-step PCR method. In the first step, two approximately 300-bp regions were amplified using genomic DNA of the WT 972 strain as a template, corresponding to the forward and reverse ends of the recombination cassette. In the second step, both modules were combined with the appropriate plasmid containing the kanamycin resistance marker (kanMX6). Transformants were selected by resistance to gentamicin (G418) and correct integrations were verified by PCR. Primer sequences used for all PCR amplifications are included in Table S6.

The gene disruption strains (TP1770 and TP1771) were constructed by replacing the target open reading frames with the kanamycin resistance marker. The pFA6a-kanMX6 plasmid [26] was used as a template for construction of replacement cassettes. The overexpression strains integrating the nmt1 promoter (TP1857, TP1855, and TP1803) were constructed using the pFA6a-kanMX6-P81nmt1, pFA6a-kanMX6-P41nmt1, and pFA6a-kanMX6-P5nmt1 plasmids [26] as templates.

Metabolomic sample preparation

Metabolomic analysis was performed as previously described [17]. Briefly, cells from cultures [40 ml/sample, 3.3×10^6 cells/ml for vegetative cells, or 10^7 cells/ml for nitrogen-starved cells, respectively] were collected by vacuum filtration and immediately quenched in 25 ml of −40°C methanol. Cells were harvested by centrifugation at −20°C and constant amounts of internal standards (10 nmol of HEPES and PIPES) were added to each sample. Cells were disrupted using a Multi-Beads Shocker (Yasui Kikai, Osaka, Japan). Proteins were removed by filtering on an Amicon Ultra 10-kDa cut-off filter (Millipore, Billerica, USA) and samples were concentrated by vacuum evaporation. Finally, each sample was re-suspended in 40 μl of 50% acetonitrile and 1 μl was used for LC-MS analysis.
LC-MS analysis

LC-MS data were obtained using a Paradigm MS4 HPLC system (Michrom Biosources, Auburn, USA) coupled to an LTQ Orbitrap mass spectrometer (Thermo Fisher Scientific, Waltham, USA). LC separation was performed on a ZIC-pHILIC column (Merck SeQuant, Umeå, Sweden; 150×2.1 mm, 5 μm particle size). Acetonitrile (A) and 10 mM ammonium carbonate buffer, pH 9.3 (B) were used as mobile phase, with gradient elution from 95% A to 20% A in 30 min at a flow rate of 100 μl/min. Peak areas of metabolites of interest were measured using the MZmine 2.10 software [51] and normalized by the weighted contribution of the areas of metabolites of interest.

Viability measurement

Cell viability was measured by plating approximately 300 cells on a YE agar plate, incubating the plate at 26°C for 4–5 days, and counting the number of colonies formed. Viability was calculated as the percentage of the number of formed colonies against the number of colonies formed at the first time point.

Supporting Information

Figure S1 Amino acid sequence alignment of S. pombe SPBC1604.01, S. japonicus SJAG_00832, N. crassa Egt-1, M. smegmatis EgtD, and M. smegmatis EgtB proteins. Alignment was generated using the COBALT algorithm. Conserved domains are indicated according to their location in S. pombe SPBC1604.01.

(TIF)

Figure S2 Amino acid sequence alignment of M. smegmatis EgtE protein and its four putative homologs in S. pombe. Alignment was generated using the COBALT algorithm. The conserved catalytic residue (PLP binding site) is indicated by a red arrow.

(TIF)

Figure S3 Normalized peak areas of EGT and its precursors obtained by metabolomic analysis of WT, the ΔSPBC660.12c single deletion mutant, and multiple deletion mutants with other putative homologs of...
mycobacterial EgtE. Cells were nitrogen-starved prior to analysis (24 h in EMM2-N medium) to induce EGT synthesis.

Figure S4  **Amino acid sequence alignment of *S. pombe* SPBC1604.01 (Egt1) protein and its closest homologs in selected species.** Alignment was generated using the COBALT algorithm. Conserved domains are indicated according to their location in *S. pombe* Egt1.

(TIF)

Figure S5  **Amino acid sequence alignment of *S. pombe* SPBC660.12c (Egt2) protein and its closest homologs in other species.** Alignment was generated using the COBALT algorithm. Conserved domains are indicated according to their location in *S. pombe* Egt2. The conserved catalytic residue (PLP binding site) is indicated by a red arrow.

(TIF)

Figure S6  **Absolute quantification of EGT content in cells.** A calibration curve was constructed by performing LC-MS injections of pure ergothioneine in 10-fold dilution steps, containing a constant amount of HEPES and PIPES standards (250 pmol each) for normalization (upper panel). Normalized peak areas were plotted against injected amounts and a regression curve was generated using Microsoft Excel. The formula to calculate absolute amount (fmol) from normalized peak area was derived from the regression curve formula (middle panel). Normalized peak areas of EGT were converted into absolute intracellular concentrations using estimated average cellular volumes (bottom panel).

(TIF)

Figure S7  **Spot test results on hydrogen peroxide and tert-butyl hydroperoxide agar plates.** WT, deletion, and overexpression strains described in this manuscript were serially diluted and grown on EMM2 plates supplemented with increasing concentrations of hydrogen peroxide (H2O2) and tert-butyl hydroperoxide (t-BOOH). The stress-sensitive Δαγ1 strain was used as a positive control.

(TIF)

Figure S8  **Cell number increase in liquid EMM2 medium supplemented with Na2SeO4.** Relative cell number increase in 24 h was measured in liquid EMM2 medium supplemented with increasing concentrations of Na2SeO4. Cell cultures were incubated at 26°C.

(TIF)

Figure S9  **Analysis of a mixture of EGT and selenium in vitro.** Extracted ion chromatograms of EGT and selenoneine masses are shown for 1 mM EGT, 1 mM Na2SeO4, and mixture of both, incubated at room temperature for 24 h. Note that the intensity scale of the selenoneine plot is 0.1% relative to that of the EGT plot.

(TIF)

Figure S10  **Verification of the identity of selenoneine and hercynylselenocysteine by their isotopic patterns.** Comparison of detected vs. calculated isotope distribution patterns of selenoneine [A] and hercynylselenocysteine [B]. Theoretical isotopic patterns were generated from the corresponding chemical formulas using the Xcalibur software (Thermo Fisher Scientific, Waltham, USA).

(TIF)

Figure S11  **Spot test results on Na2SeO4 agar plates.** WT, deletion, and overexpression strains described in this manuscript were serially diluted and grown on EMM2 plates supplemented with increasing concentrations of Na2SeO4. The stress-sensitive Δαγ1 strain was used as a positive control.

(TIF)

Table S1  **Locations and amino acid sequences of conserved protein domains shown in Figure 1B and Figure S1.**

(XLSX)

Table S2  **Locations and amino acid sequences of conserved protein domains shown in Figure 1C and Figure S2.**

(XLSX)

Table S3  **Results of all LC-MS measurements of the intermediates in the EGT biosynthetic pathway.**

(XLSX)

Table S4  **Locations and amino acid sequences of conserved protein domains of Egt1 homologs shown in Table 2 and Figure S4.**

(XLSX)

Table S5  **Locations and amino acid sequences of conserved protein domains of Egt2 homologs shown in Table 2 and Figure S5.**

(XLSX)

Table S6  **Sequences of oligonucleotide primers used for PCR amplifications.**

(XLSX)

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**Author Contributions**

Conceived and designed the experiments: TP. Performed the experiments: TP. Analyzed the data: TP. Contributed reagents/materials/analysis tools: TP MU. Wrote the paper: TP MY. Provided supervision and guidance: TP.

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