Lack of mycothiol and ergothioneine induces different protective mechanisms in *Mycobacterium smegmatis*

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**Abstract**

*Mycobacterium smegmatis* contains the low molecular weight thiols, mycothiol (MSH) and ergothioneine (ESH). Examination of transposon mutants disrupted in *mshC* and *egtA*, involved in the biosynthesis of MSH and ESH respectively, demonstrated that both mutants were sensitive to oxidative, alkylating, and metal stress. However, the *mshC* mutant exhibited significantly more protein carbonylation and lipid peroxidation than wildtype, while the *egtA* mutant had less protein and lipid damage than wildtype. We further show that Ohr, KatN, and AhpC, involved in protection against oxidative stress, are upregulated in the *egtA* mutant. In the *mshC* mutant, an Usp and a putative thiol peroxidase are upregulated. In addition, mutants lacking MSH also contained higher levels of Coenzyme F420 as compared to wildtype and two Coenzyme F420 dependent enzymes were found to be upregulated. These results indicate that lack of MSH and ESH result in induction of different mechanisms for protecting against oxidative stress.

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**1. Introduction**

Mycothiol (MSH), the major low-molecular weight (LMW) thiol produced by Actinomycetes, acts as a glutathione (GSH) analog to maintain redox homeostasis, and protects the cell against xenobiotic agents, notably in pathogens such as *Mycobacterium tuberculosis* [1,2]. MSH is present in mycobacteria at millimolar levels [1] and is synthesized in a five-step pathway from glucose-6-phosphate with the enzymes MshA, A2, B, C, D [2]. A second LMW thiol, l-ergothioneine (ESH), is also synthesized by Actinomycetes but is present in lesser amounts [3]. In *M. smegmatis*, ESH biosynthesis is initiated by a methyltransferase (EgtD), and proceeds through an additional five-step pathway involving the enzymes EgtA-E [4]. Unlike other LMW thiols as MSH, ESH is tautomeric [5] and exists predominantly in the thione form in neutral aqueous solutions. In addition, the redox potential of ESH is −60 mV as compared to −250 mV of GSH which may account for ESH’s resistance to auto-oxidation [5].

While the role and function of MSH has been extensively studied, the role of ESH in bacteria is not as well-described. MSH mutants are more susceptible than wildtype to a number of antibiotics, oxidative stress, alkylating agents, and other stresses but resistant to the pro-drugs, isoniazid and ethionamide [6]. An *M. smegmatis* mutant disrupted in *egtD*, which lacks ESH, and is more susceptible to the lipid peroxides, cumene hydroperoxide (CHP) and tert-butyl hydroperoxide (tBOH), but not sensitive to the antibiotics, rifampin, isoniazid, ethionamide, and ethambutol [3]. A double mutant disrupted in *mshA* and *egtD* lacking both MSH and ESH is significantly more sensitive to the peroxides than either of the single mutants lacking either ESH or MSH. Interestingly, levels of ESH are significantly higher in the *mshA* mutant indicating that ESH may be able to compensate for the loss of MSH [7]. In addition, Ohr, an organic hydroperoxide reductase, is substantially upregulated in both *mshA* and *egtD* mutants. Whether this upregulation of Ohr is particular to *mshA* and *egtD* mutants or is common in mutants disrupted in other genes in the biosynthetic pathway of MSH and ESH is unknown.

In a screen for genes involved in disulfide stress, a transposon mutant library was screened for sensitivity to diamide, a thiol-oxidizing agent. Five mutants lacking MSH and disrupted in *mshC*, *S24*, *D43*, *D48*, *D41*, and *R41*, and a mutant disrupted in *egtA*, *R119*, were identified as having increased sensitivity to diamide. We demonstrate that the *mshC* mutant, *S24*, has significantly more protein carbonylation and lipid peroxidation, measures of oxidative damage, while the *egtA* mutant has less protein and lipid damage than wildtype. We further show that Ohr and other enzymes involved in protection against oxidative stress are

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upregulated in the egtA mutant, while in the mshC mutant, the levels of coenzyme F420, a coenzyme involved in redox reactions, are increased.

2. Materials and methods

2.1. Bacterial strains, culture conditions, and disk sensitivity assays

*M. smegmatis* mc²155 and mutant strains were grown in Middlebrook 7H9 broth (Difco) with 10% OADC and then diluted in Middlebrook 7H9 broth with 0.05% Tween 80% and 1% glucose. The strains were also grown on solid medium with 0.5% glycerol with 1% glucose for disk sensitivity assays, which were performed as previously described [6]. Antibiotics were added to the media for the transposon mutant (Kanamycin 25 μg/ml) and complemented strains (Hygromycin 75 μg/ml). Broth cultures were grown in a shaking incubator and all cultures were grown at 37 °C unless otherwise indicated.

2.2. Creation and screening of a transposon mutant library and identification of the disrupted gene

An *M. smegmatis* transposon mutant library (EZ::TN < kan-2 > Tnp transposase and Tn5 kanamycin resistance marker) was constructed and screened for diamide sensitive mutants as described by Rawat et al. [8]. To identify the site of insertion, genomic DNA was digested with restriction enzymes SaII and PstI followed by self-ligation and PCR amplification with primers complementary to the transposon. The sequence of the amplified PCR product was compared to the *M. smegmatis* genome sequence [8].

2.3. Determination of thiol levels and Coenzyme F420

Thiols in cell extracts were labeled with monobromobimane (mBBr) as previously described [6]. High-pressure liquid chromatographic (HPLC) analysis was carried out on a HiChrom Ultra-sphere Ion Pair column (5 μm; 250 × 4.6 mm) with the following gradient at 1 ml/min of Buffer A, 0.25% glacial acetic acid, pH 4.0; and Buffer B, methanol (0–5 min, 10% A; 35 min, 18% A; 45 min 27% A; 47 min, 100% A; 49 min, 0%). HPLC analysis for F420 content was performed as previously described [9] with the following modifications: Shimadzu Prominence system (LC 20 CE dual pumps, SIL-20A autosampler and SPD-M20A Diodearray detector) was used for chromatographic separation and quantitative analysis. Cell extracts were separated with a Vydac 218TP54 column (C18, 5 μm, 4.6 mm i.d. × 250 mm) at a flow rate of 0.6 ml/min. A linear gradient of sodium acetate buffer-acetonitrile solution was used and elution was monitored at 400 nm. Identities of F420 peaks were confirmed by their UV-Visible spectrum collected by an in-line diode array detector. Samples in quadruplicates were analyzed at least three times and reported as averages ± SD. Student *t* test was performed with *p* ≤ 0.05 regarded as significant.

2.4. Complementation of S24 (mshC mutant), R119 (egtA mutant)

*M. tuberculosis* mshC cloned into pSODIT [10] was introduced into S24 in order to obtain the complemented strain, S24C. Primers, TeshA1, with a HindIII restriction site (CTTAAAGCTTCCAC-GACGAGGCCTGGAC) and TeshA2 with a Xhol restriction site (TGCTCTAGGTAAGTCTAGGACGCCCG) were used to amplify *M. tuberculosis* egtA (Rv3704c) and 297 base pairs upstream of the start codon, which contained the promoter region of the gene. The 1609 bp amplicon and complementation vector pHINT were digested with the HindIII and Xhol, ligated and transformed into *E. coli*. After confirmation of cloning, the resulting vector, pHINTegtA, was electroporated into R119 competent cells. Kanamycin and hygromycin resistant transformants were screened by PCR for *M. tuberculosis* egtA and one colony, R119C, was characterized further.

2.5. Measurement of protein carbonylation, lipid peroxidation, and peroxidase activity

Protein carbonylation and lipid peroxidation was measured using the Protein Oxidation Detection Kit and TBARS Assay Kit, following the manufacturer’s instructions (Cayman Chemical). To assess protein carbonylation, cells were grown until OD₆₀₀ equaled one. Then one set of 10 ml cultures in triplicate were treated with 5 mM H₂O₂ and another set was left untreated for one hour. The cultures were harvested and lysed using bead beating (Fast Prep FP120, Thermo Electron company). The cell lysates were centrifuged at 13,000 rpm for 30 min at 4 °C to get rid of cell debris and the protein concentration of the supernatant was determined by Bio-Rad DC assay. Two mg/ml of protein was incubated with 2,4-dinitrophenylhydrazine at room temperature in the dark for one hour with frequent vortexing. The derivatized proteins were TCA precipitated and the pellet was further extracted with ethanol/ethanol acetate (1:1) solution three times. The resulting pellet was resuspended in guanidine hydrochloride. After centrifugation, the supernatant containing the protein hydrazones were analyzed spectrophotometrically at 360 nm.

Lipid peroxidation was assessed by treating 1 ml of cells in triplicate at 1.0 OD₆₀₀ in log phase with 30 mM FeSO₄ and 5 mM H₂O₂ for two hours. Cell lysates were prepared similarly to the samples for protein carbonylation assay and to 100 μl of the cell lysate, 100 μl of 10% SDS was added. A color reagent was prepared by adding thiobarbituric acid (TBA) to acetic acid followed by the addition of sodium hydroxide to the solution. Four ml of the color reagent was added to each sample and the samples were boiled for 1 h and then transferred to ice for 10 min to stop the reaction. After centrifugation at 1600 x g for 10 min at 4 °C, the supernatants were transferred to an ELISA plate and A₅₃₂ was determined in duplicate. The amount of lipid peroxidation was calculated with reference to a standard curve of malondialdehyde.

Peroxidase activity was determined for each strain in triplicate using the Pierce Quantitative Peroxide Assay Kit, which detects levels of peroxides based on the oxidation of ferrous to ferric ion at acidic pH. The cell free lysate was prepared as described for the protein carbonylation, 300 μg of protein was incubated with 45 μM H₂O₂, CHP, and TbOH separately for 30 min. The amount of remaining peroxides was determined by addition of sulfuric acid solution containing ferrous ion and xylene orange and incubation at room temperature for 20 min. The ferric ion-xylene orange purple complex was measured at 595 nm. The amount of peroxides was calculated with reference to a standard curve of the relevant peroxide.

All the experiments were performed at least three times.

2.6. Proteomic evaluation of wildtype and mutants, S24 and R119

As described in Ta et al. [7], *M. smegmatis* strains were grown for 11 days on Middlebrook 7H9 agar and 1.0% glucose at 23 °C. Cells were collected by centrifugation and resuspended in lysis buffer (50 mM Tris containing 7 M urea, 2 M thiourea, and 4% 3-[(3-Cholamidopropyl) dimethylammonio]-1-propanesulfonate (CHAPS)). Two-dimensional difference gel electrophoresis (2D-DIGE) analysis of the supernatant was performed at Applied Biosystems (Hayward, CA). Briefly, samples were fluorescently labeled and subjected to isoelectric focusing (IEF) on a 13 cm pH gradient strip (pH 4–6). The changes in abundance of proteins between the wildtype and the mutants were identified using the ImageQuant TL and DeCyder softwares. Spots were excised from the gel and

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identified by MALDI TOF/TOF (tandem mass spectrometry MS/MS) on a 5800 mass spectrometer (AB Sciex). Resulting peptide masses and associated fragmentation spectra were submitted to GPS Explorer version 3.5 with the MASCOT search algorithm (Matrix science) and searched against the National Center for Biotechnology Information non-redundant database. Searches were performed without constraining protein molecular weight or isoelectric point, with variable carboxamidomethylation of cysteine and oxidation of methionine residues, and with one missed cleavage allowed in the search parameters. MASCOT searches were also performed to identify phosphorylation in the proteins. Candidates with score of either protein C.I.% or Ion C.I.% greater than 95 were considered significant.

3. Results

3.1. Mycothiol, ergothioneine and coenzyme F420 levels in transposon mutants lacking MSH and ESH

Inverse PCR was performed to identify the disrupted genes in diamide sensitive mutants, S24, D41, D43, D48, R41, and R119. Five of the mutants, S24, D41, D43, D48, R41, are disrupted in msfC (Msmeg.4189) and did not contain any MSH. In four of the five mutants disrupted in msfC, the transposon was inserted at 589 nucleotides from the start codon. In the D48 mutant, the insertion site was 599 nucleotides from the start codon. As the majority of the mutants had the same transposon insertion site, we decided to focus on only one of these mutants, S24, further. The amount of ESH in S24, 0.37 ± 0.04 μmol/g dry cell weight, was higher than in wildtype (0.20 ± 0.01) and significantly different. The increase in ESH was not as high as in the mshA mutant (1.02 ± 0.24) grown under the same growth conditions [7]. The complemented strain, S24C, did not show a significant difference from wildtype in ESH levels (0.19 ± 0.01) and MSH levels (wildtype: 6.21 ± 0.17, S24C: 5.90 ± 0.66).

In the transposon mutant, R119, where the transposon is inserted 972 nucleotides after the start codon in egtA (Msmeg.6250), there was no ESH present and MSH levels (6.10 ± 0.37) were not statistically significantly different than wildtype. The complemented strain, R119C, did not have statistically significant different levels of MSH (5.88 ± 0.27) and ESH from wildtype (0.24 ± 0.04).

Previously, we had demonstrated that mutants disrupted in fiHC, which is involved in the biosynthesis of Coenzyme F420 (F420), a deazaflavin derivative of FMN found in methanogenic Archaea and actinobacteria [11], are sensitive to oxidative stress [12,13]. We measured F420 levels to determine if F420 can compensate for the lack of MSH or ESH. F420 levels were statistically significantly higher in mshA (8.47 ± 0.52 nmol/g wet cell weight) and msfC (S24) mutants (8.05 ± 0.24) as compared to wildtype (6.12 ± 0.87). Complementation of the mshA mutant showed partial restoration of wildtype F420 levels (7.16 ± 0.57) while complementation of S24 resulted in lower F420 levels (4.81 ± 0.32) as compared to wildtype. Interestingly, R119 had statistically significantly less F420 (4.21 ± 0.11) than the wildtype and complementation (6.51 ± 0.38) restored F420 to wildtype levels.

3.2. Sensitivity of S24 and R119

In liquid culture, there is no difference in growth between R119 and wildtype under normal growth conditions (data not shown). However, R119, like S24 is more susceptible to alkylating agents and the sensitivity is reversed by complementation with the M. tuberculosis homolog (Table 1). In the case of oxidants, both mutants show sensitivity to diamide and CHP, but only S24 demonstrates sensitivity to H2O2.

As metal stress results in oxidative stress through either the Fenton reaction or sequestration of LMW thiols, we also examined metal sensitivity. Both mutants were sensitive to Cd2+, Cu2+, SeO3−, TeO3−, and CrO42−, but only S24 was more sensitive to Fe2+, as compared to wildtype (Table 1). The sensitivity to oxidants, alkylating agents, and metals was reversed in R119C and partially reversed in S24C.

3.3. Protein carbonylation and lipid oxidation in mutants lacking MSH and ESH

Protein carbonylation, as an indicator for protein damage, was measured in S24 and R119, with and without hydrogen peroxide. S24 demonstrated significantly higher levels of protein carbonylation which was partially reversed in the complemented strain, S24C. In contrast, R119 had lower levels of protein carbonylation and the complemented strain, R119C, was not significantly different from the wildtype. S24 also had significantly higher levels of lipid peroxidation and R119 had lower lipid oxidation levels. The complemented strain, R119C, was not significantly different from wildtype (Table 2). Thus, lack of MSH results in increased damage while lack of ESH results in lesser damage to cellular proteins and lipids.

3.4. Peroxidase activity in mutants lacking in MSH and ESH

To determine if MSH or ESH are involved as electron donors to a peroxidase(s), the decomposition of H2O2, CHP, and tBOH was measured using xylene orange assay. The amount of peroxides was higher in S24 than wildtype (Table 2) indicating that MSH is able to either directly reduce the peroxides or act as an electron donor to a peroxidase. In contrast, the amount of peroxides in R119 was the same as wildtype (Table 2) indicating that ESH is not directly involved in the reduction of peroxides or another mechanism is induced in R119 that detoxifies peroxides. The peroxide levels in S24C were similar to wildtype.

| Table 1 | Susceptibility of Mycobacterium smegmatis ergothioneine mutants to antibiotics, toxins, metals and oxidants as determined by disk assays. All values represent averages and SD (n=4). Student t-test was performed and the p values of mutants were compared with the wildtype. |
|---|---|
| **Reagent** | **Zone of clearing (mm)** A.R. Singh et al. / Biochemistry and Biophysics Reports 8 (2016) 100–106 |
| **Oxidants** | Wt | S24 | S24C | R119 | R119C |
| H2O2 (0.5 μmol) | 20 ± 1 | 26 ± 2* | 22 ± 2* | 22 ± 1 | 19 ± 1 |
| Diamide (15 μmol) | 19 ± 1 | 27 ± 1** | 25 ± 1** | 29 ± 1** | 20 ± 2 |
| CHP (0.5 μmol) | 23 ± 2 | 31 ± 2** | 27 ± 1 | 29 ± 1** | 27 ± 2* |
| Alkylating agents |  |
| N-ethylmalamide (0.25 μmol) | 14 ± 2 | 23 ± 1** | 16 ± 1 | 22 ± 1** | 15 ± 1 |
| Iodoacetamide (0.0125 μmol) | 16 ± 3 | 21 ± 2* | 18 ± 2 | 22 ± 2* | 19 ± 2 |
| Chlorodinitrobenzene (0.0125 μmol) | 17 ± 2 | 22 ± 1** | 18 ± 1 | 21 ± 1* | 18 ± 2 |
| **Metal** |  |
| Cd2+ (0.5 μmol) | 22 ± 2 | 31 ± 1** | 27 ± 1* | 27 ± 1* | 23 ± 1 |
| Cu2+ (0.5 μmol) | 19 ± 1 | 22 ± 2* | 20 ± 1 | 22 ± 2* | 18 ± 2 |
| SeO3− (0.25 μmol) | 20 ± 1 | 25 ± 1** | 19 ± 1* | 23 ± 1** | 22 ± 2 |
| TeO3− (0.25 μmol) | 15 ± 2 | 22 ± 1** | 18 ± 1* | 19 ± 1** | 16 ± 1 |
| Fe2+ (5 μmol) | 16 ± 1 | 21 ± 3* | 17 ± 3 | 19 ± 4 | 17 ± 3 |
| CrO42− (0.5 μmol) | 20 ± 1 | 27 ± 2** | 24 ± 1* | 24 ± 1** | 22 ± 1 |

* *p < 0.0005 using student’s t-test.
** p < 0.05 using student’s t-test.
*** p < 0.005 using student’s t-test.
Table 2

| Strains | Protein carbonylation (μM L⁻¹) | Lipid oxidation (μM) | Peroxidase activity (μM) |
|---------|--------------------------------|---------------------|-------------------------|
|         | Untreated | 5 mM H₂O₂ | Untreated | 5 mM H₂O₂ | CHP | H₂O₂ | TBOH |
| WT      | 15.4 ± 2.3 | 26.9 ± 4.6 | 201.1 ± 3.3 | 15.4 ± 2.3 | 26.9 ± 4.6 | 504.0 ± 4.3 | 75.0 ± 2.9 | 20.1 ± 0.4 | 15.4 ± 2.3 | 26.9 ± 4.6 | 70 ± 0.7 |
| S24     | 36.2 ± 3.9*** | 55.8 ± 2.1*** | 90.0 ± 1.9*** | 102.4 ± 2.4*** | 35.0 ± 1.9*** | 33.0 ± 0.7*** | 24.5 ± 0.8*** |
| S24C    | 29.1 ± 2.4*  | 40 ± 3.1***  | 79.1 ± 3.8 | 19.0 ± 0.6 | 17.5 ± 2.0 | 15.0 ± 0.9 |
| R119    | 8.9 ± 1.9**  | 21.2 ± 4.1†  | 43.3 ± 1.3*  | 57.1 ± 5.1**  | 18.1 ± 1.6 | 16.8 ± 1.1 | 17.6 ± 1.0 |
| R119C   | 15.0 ± 3.3  | 26.0 ± 4.9  | 39.0 ± 4.9  | 62.6 ± 5.0  | 19.0 ± 1.2 | 18.5 ± 1.0 | 19.0 ± 1.9 |

Table 2: Protein carbonylation, lipid peroxidation, and peroxidase activity in M. smegmatis mutants and complemented strains. All values represent averages and SD (n=3). Student t-test was performed and the p values of mutants were compared with the wildtype.

3.5. 2D-DIGE analysis of S24

Since standard SDS-PAGE analysis of random mutations disrupted in mshA and egtD demonstrated the overproduction of a band identified as Ohr [3,7], 2D-DIGE was performed on wildtype and S24. In S24, 36 spots were clearly different between S24 and wildtype and of these 19 were conclusively identified (Table 3).

Table 3

| Spot | S24/wt | M. smegmatis | M. tuberculosis | Description |
|------|--------|--------------|-----------------|-------------|
| 18   | 8.12   | MSMEG_4283   | Rv2215          | SycB/Dlt4 F420 reduc-tase, dihydrodolipoyl-oxidoreductase |
| 14   | 4.5    | MSMEG_4688   | Rv2466c         | Thiol peroxidase, dihydrolipoamide dehydrogenase |
| 11   | 8.8    | MSMEG_2027   | Rv1436          | Glyceraldehyde-3-phosphate dehydrogenase |
| 24   | 7.63   | MSMEG_1996   | Rv579c          | Coenzyme F420 dependent peroxidase and peroxynitrite reductase |
| 16   | 5.17   | MSMEG_0415   | Rv2026c         | Putative Coenzyme F420 reductase, NADH-FMN oxidoreductase |
| 23   | 5.98   | MSMEG_3432   | Rv2158          | NADH dehydrogenase, FAD oxidoreductase |
| 19   | 8.36   | MSMEG_4362   | Rv2158          | NADH dehydrogenase, FAD oxidoreductase |
| 22   | 6.76   | MSMEG_3811   | Rv1636          | Usp oxidoreductase |
| 27   | 4.55   | MSMEG_3811   | Rv1636          | Usp oxidoreductase |
| 28   | 5.08   | MSMEG_3811   | Rv1636          | Usp oxidoreductase |
| 29   | 5.28   | MSMEG_3811   | Rv1636          | Usp oxidoreductase |
| 13   | 5.13   | MSMEG_0880   | Rv0440          | GroEL chaperonin |
| 26   | 37.25  | MSMEG_6431   | Rv3849          | EspR nitrogen regulatory protein |
| 30   | 4.83   | MSMEG_2426   | Rv2919c         | GlnA nitrogen regulatory protein |
| 1    | –3.12  | MSMEG_2426   | Rv2220          | GluA synthetase, SycB |
| 9    | 2.59   | MSMEG_2426   | Rv2890c         | RpsB |
| 13   | 12.29  | MSMEG_1401   | Rv0695 Tuf |
| 35   | 12.67  | MSMEG_1401   | Rv0695 Tuf |
| 8    | 3.03   | MSMEG_1401   | Rv0695 Tuf |
| 15   | 3.69   | MSMEG_5050   | No homolog      | Map |

**P ≤ 0.0005 using student’s t-test.**
***P ≤ 0.005 using student’s t-test.**
*P ≤ 0.05 using student’s t-test.

Supp Fig. 1). A number of proteins associated with protection against oxidative stress are up-regulated in S24. One of these upregulated proteins is MSMEG_4283 (Spot 18), dihydrodolipoyl-oxidoreductase (Dlt4), which is the E2 component of pyruvate dehydrogenase complex, and contains a lipoyl binding site [14]. M. tuberculosis dlat mutant has severe growth defect in vitro, which has reduced survival in bone marrow-derived macrophages in c57b/6 and inos –/– mice, and is less able to grow in c57b/6 mouse lungs, spleen and liver in vivo [15]. Together with lipoyamido dehydrogenase (LpdC), AhpD, and AhpC, DlaT acts as an NADH dependent peroxidase and peroxynitrite reductase to provide protection against oxidative stress [14]. The importance of this dihydrolipoyldehydrogenase system is highlighted by the discovery of suppressor mutations in lipoyamido dehydrogenase (LpdA) in E. coli. An E. coli mutant lacking both glutathione reductase and thioredoxin reductase is not viable but suppressor mutations in lpdA restore growth to this redox-defective mutant by causing the accumulation of dihydrolipoyldehydrogenase, which reduces oxidized glutaredoxins and circumvents the need for thioredoxin and glutathione in E. coli [16].

MSMEG_4688, also upregulated, codes for a putative thiol peroxidase and its M. tuberculosis ortholog, Rv2466c, has been shown to be involved in activation of TIP053, a thienopyrimidine compound considered to be a leading drug target against tuberculosis [17]. Rv2466c is a member of the sigH regulon and is induced in the Wayne model of dormancy [18]. The physiological electron donor for this thiol peroxidase is unknown although it is able to reduce peroxides using DTT as an electron donor [17]. Interestingly, two of the proteins up-regulated include coenzyme F420 dependent proteins: MSMEG_1996 (Spot 16), a methylthiazoyldihydrocarboxanetin reductase that reduces oxidized coenzyme F420, and MSMEG_2027 (Spot 24), a quinone reductase, which uses F420 as an electron donor or repressor to activate another new tuberculosis drug, PAS-824 [13]. Another indication that S24 is under oxidative stress is the upregulation of GAPDH, a marker of oxidative stress, which is present in two spots (11 and 12) that...
represent different post-translational modifications in the form of oxidation cysteine residues to sulfenic, sulfonic and sulfenic acid.

Several other proteins that are upregulated in S24 are associated with a general stress response. Two Usps (universal stress proteins), MSMEG_4362 (Spot 19) and MSMEG_3811 (Spots 22, 27, 28, 29), are differentially expressed. MSMEG_3811 codes for an iron regulated single domain Usp and is the only Usp which has a homolog in M. leprae. The homolog in M. tuberculosis, Rv1636, is the only Usp along with kdpD, which is not up-regulated during macrophage infection [19]. Of the spots representing MSMEG_3811, spots 22 and 28 are up-regulated in the wildtype, and spots 27 and 29 are up-regulated in S24. A MASCOT search for phosphorylations revealed that the wildtype MSMEG_3811 isoforms are phosphorylated while the msfC isoforms are not.

Spot 13 is the 65 kDa heat shock protein, GroEL (MSMEG_0880), which prevents misfolding and promotes the refolding and proper assembly of unfolded polypeptides generated under stress conditions like high temperatures. Spot 26 is MSMEG_6431, the homolog of M. tuberculosis EspR (Rv3849), which regulates ESX-1, required for secretion and virulence in mice [20]. RpsB (Spot 33), 30s ribosomal protein S2, is involved in formation of the translation initiation complex and involved in the metabolism of ppGpp, which mediates the stringent response that coordinates a variety of cellular activities in response to changes in nutritional abundance. EspR, Tuf (Spot 8, 35) and the Usp, MSMEG_3811, are regulated by iron levels, suggesting an interaction among iron levels, oxidative stress, and LMW thiols.

Also upregulated in the mutant is nitrogen regulatory protein PII (Spot 30), which becomes uridylylated to pII-UMP by GlnD when the ratio of glutamine to 2-ketoglutarate decreases. The pII-UMP allows the deadenylylation of glutamine synthetase activating enzyme. During nitrogen excess, p-II is deuridylated and UMP allows the deadenylylation of glutamine synthetase activity. PII (Spot 30), which becomes uridylylated to pII-UMP by GlnD, is involved in formation of the translation initiation complex and involved in the metabolism of ppGpp.

3.6. 2D-DIGE analysis of R119

The DIGE protein profile of wildtype and R119 demonstrated that 98 proteins were differentially regulated (Table 4, Supp Fig. 2). Of these 17 spots had an egtaA/wt ratio greater than two and 18 spots had an egtaA/wt ratio less than –2. Ohr is the most upregulated protein and was present in four spots (Spots 88, 89, 90, 91).

Oxidation of specific cysteines and perhaps mycothiolation of other cysteines may be responsible for these multiple spots of Ohr. In addition to Ohr, other proteins associated with protection against oxidative stress, AhpC (Spot 72), another thiol peroxidase, and KatN (Spot 48), a manganese containing catalase, are also up-regulated in R119. Another up-regulated protein, MSMEG_4272 (Spot 95), is a member of an iron-sulfur cluster biosynthesis protein family, which have been shown to be dependent on LMW thiols such as bacillithiol for assembly [21]. In addition, a MarK type regulator (Spot 82), upstream of a quinone reductase, is upregulated.

Proteins associated with metabolism are differentially expressed in R119 and wildtype. Lactate monooxygenase (Spot 36), which catalyzes the conversion of lactate to acetate, is upregulated while glycerol kinase (Spot 22), which catalyzes the transfer of a phosphate from ATP to glycerol dihydroxyacetone, l-glycer-aldehyde and D-glycer-aldehyde during carbohydrate metabolism, is the most downregulated protein. IlvC (Spot 40) associated with isoleucine and valine metabolism is also downregulated. Other downregulated proteins include an alcohol dehydrogenase (Spot 44), belonging to family of quinone reductases, a flavin-containing monooxygenase (Spot 33), and MPT63 (Spot 19), an immunogenic protein which has been implicated in virulence.

In contrast to S24, the Usp, MSMEG_3811, is down-regulated in R119. GroEL, another general stress protein, is also down-regulated in R119 but up-regulated in S24. Other protein markers for stress seen in S24 are not up-regulated in R119 and R119, thus, does not appear to be in a “stressed” state. 2D-DIGE of R119C demonstrated partial complementation by the M. tuberculosis egtaA gene. In particular, the R119C/wt ratios for spots representing Ohr are significantly lower as compared to R119/wt. Although other proteins are differentially expressed in both R119 and R119C, the R119C/wt ratios are less than R119/wt ratios. As other phenotypic data for S24 demonstrated that this strain was partially complemented by the M. tuberculosis msfC, 2D-DIGE was not performed.

4. Discussion

Herein, we confirm that both MSH and ESH protect mycobacterial cells against a number of stresses, including oxidative stress and alkylating stress, and demonstrate that these two LMW proteins associated with metabolism are differentially expressed in R119 and wildtype. Lactate monooxygenase (Spot 36), which catalyzes the conversion of lactate to acetate, is upregulated while glycerol kinase (Spot 22), which catalyzes the transfer of a phosphate from ATP to glycerol dihydroxyacetone, l-glycer-aldehyde and D-glycer-aldehyde during carbohydrate metabolism, is the most downregulated protein. IlvC (Spot 40) associated with isoleucine and valine metabolism is also downregulated. Other downregulated proteins include an alcohol dehydrogenase (Spot 44), belonging to family of quinone reductases, a flavin-containing monooxygenase (Spot 33), and MPT63 (Spot 19), an immunogenic protein which has been implicated in virulence.

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Table 4

Differentially expressed proteins in wildtype, egtaA mutant, R119, and complemented strain, R119C.

| Spot | R119/WT | R119C/WT | M. smegmatis | M. tuberculosis | Description |
|------|---------|----------|--------------|-----------------|-------------|
| 89   | 254.25  | 1.48     | MSMEG_0447   | –               | Ohr, Organic hydroperoxide protein |
| 88   | 75.47   | 1.19     | MSMEG_0447   | –               | –            |
| 90   | 18.4    | 1.18     | MSMEG_0447   | Rv2204c         | HesB/YadR/Yvh family protein | iron sulfur proteins |
| 91   | 8.6     | –2.8     | MSMEG_0447   | –               | Lactate-2-monooxygenase |
| 95   | 21.51   | 14.85    | MSMEG_4272   | Rv4248          | AhpC alkylhydroperoxide reductase |
| 36   | 10.46   | 26.52    | MSMEG_3962   | Rv1636          | MarR, transcriptional regulator, upstream of quinone reductase |
| 72   | 8.58    | 6.61     | MSMEG_4891   | Rv1926c         | MspA, Chain A, Rim Domain Of Main Porin |
| 82   | 5.36    | 6.82     | MSMEG_6361   | Rv3001c         | KatN, manganese containing catalase |
| 78   | 3.62    | 6.47     | MSMEG_0965   | Rv4040          | Aminoglycoside phosphotransferase-kanamycin resistance gene |
| 83   | 3.45    | 2.75     | MSMEG_6213   | Rv1311          | – Conserved hypothetical protein |
| 52   | 3.34    | 1.81     | MSMEG_1680   | GroEL            | – Universal stress protein family protein |
| 100  | 2.34    | 2.11     | MSMEG_1680   | AtpC, FOF1 ATP synthase subunit epsilon |
| 101  | –1.07   | –1.39    | MSMEG_3811   | Rv1636          | Universal stress protein family protein |
| 102  | –1.8    | –1.45    | MSMEG_4935   | Rv6640          | GroEL |
| 10   | –1.87   | 1.19     | MSMEG_0880   | –               | Protocatechuate 3,4-dioxygenase/acetyl-CoA thioesterase II |
| 59   | –3.24   | –1.45    | MSMEG_2601   | –               | – Protocatechuate 3,4-dioxygenase/acetyl-CoA thioesterase II |
| 40   | –4.66   | –2.12    | MSMEG_2374   | Rv3001c         | IlvC, Keto-acid isomerase |
| 15   | –5.44   | 3.85     | MSMEG_4303   | –               | Methyltransferase |
| 19   | –5.7    | –2.71    | MSMEG_3412   | Rv1926c         | Immunogenic protein MPT63 |
| 33   | –6.33   | –4.28    | MSMEG_1682   | –               | Flavin-containing monooxygenase FMO |
| 44   | –7.76   | –6.65    | MSMEG_2079   | –               | Alcohol dehydrogenase, Yhd/Yhp family |
| 22   | –8.24   | –9.5     | MSMEG_6759   | Rv3696c         | GlpK, Glyceraldehyde kinase |
thiols are also involved in protection against metal stress. S24 is extremely sensitive to oxidants and alkylating agents, as previously reported for other mutants lacking MSH [6], while R119 is less sensitive (Table 1). Similar to the M. smegmatis mutant disrupted in egT, which is more susceptible to the lipid peroxides, CHP and tert-butyl hydroperoxide (tBOH), R119 is sensitive to CHP [3]. In contrast to the Streptomyces coelicolor egT mutant, R119 is not sensitive to hydrogen peroxide [22].

Both mutants are also sensitive to metal stress. In vitro studies had previously demonstrated that ESH is able to scavenge free radicals and metal ions. In particular, ESH is known to form complexes with copper [23] in vitro but in vivo, copper sulfate had no effect on conidial germination or hyphal growth in a mutant of Neurospora crassa lacking ESH [24]. Other LMW thiols like GSH are known to protect against cadmium toxicity in E. coli [25] and mutants disrupted in bacillithiol, which is structurally similar to MSH, are sensitive to metal stress [26].

Recently, Servillo et al. [27] reported that oxidation of ESH with hypochlorite, peroxynitrite and hydrogen peroxide resulted in the sulfonated form (ESO–H), and hercynine, the desulfurated form of ESH, indicating that ESH, like cysteine and MSH, is oxidized irreversibly during oxidative stress. Moreover, Sao-Emani et al. (2013) reported that a double mutant lacking both ESH and MSH is significantly more sensitive to the peroxides than either of the single mutants lacking either ESH or MSH. Taken together, these results suggest that MSH and ESH may be able to partly compensate for the loss of the other thiol. However, as the levels of ESH are 25–50 fold lower than MSH in M. smegmatis, ESH is less likely to be able to compensate for MSH. In the mshA mutant, which lacks MSH, ESH levels are substantially higher but that is not the case in the mshC mutant. Instead, the levels of Coenzyme F420 are higher in S24. As disruption of mshA and eshD results in a dramatic overproduction of organic hydroperoxide resistance protein, Ohr [3,7], we sought to establish if the same was true for mutants disrupted in mshC (S24) and eshA (R119). In S24, there is no upregulation of Ohr; however, the upregulation of DiaT which is part of the system that provides reducing equivalents to the thiol peroxidases, AhpC and Ohr, and the upregulation of the M. smegmatis homolog of the putative thiol peroxidase, Rv2466c, implies that there is a major increase in oxidative stress in the mshC mutant. Upregulation of an enzyme that likely serves as a F420-dependent quinone reductase, preventing the formation of cytotoxic semiquinones, and upregulation of a putative F420-dependent reductase implies that coenzyme F420, which has been shown to be involved in protection against oxidative stress [12], may be able to partially substitute for MSH in the mycobacterial cell.

Interestingly, multiple spots representing Usp (MSMEG_3811) are present in the 2D-DIGE. These multiple spots are a result of different amounts of phosphorylation of MSMEG_3811. Intriguingly, Usp is not phosphorylated in S24, implying a cross-talk between MSH and universal stress proteins. Previously, we observed phosphorylation of another Usps, MSMEG_3940, during stationary phase in wildtype M. smegmatis using a stain for phosphoproteins (data not shown). Phosphorylation of Usps has also been demonstrated in Arabidopsis challenged with biotic stress [28] although how this signal is translated into a protective response is not clear. Lack of MSH alters the redox environment of the cell which may cause oxidative modifications of kinases, either inhibiting or activating them [29]. The resulting phosphorylation/dephosphorylation events would positively regulate some and negatively regulate other signal transduction pathways [29]. In M. smegmatis wildtype, there would be presumably be less mycothiolation of proteins, including that of a kinase that phosphorylates Usp or perhaps less mycothiolation of MSMEG_3811 directly. This could result in phosphorylation or autophosphorylation of MSMEG_3811, as in E. coli UspG [31]. In S24, mycothiolation of the kinase would reduce activity and less phosphorylation or autophosphorylation. It is not clear what the exact functions of Usps are but it is likely that post-translation modifications of MSMEG_3811 in S24 have far-reaching consequences involving changes in metabolism and growth arrest.

Despite upregulation of genes involved in the stress response, S24 has increased protein carbonylation and lipid peroxidation. Paradoxically, R119 has less protein carbonylation and lipid oxidation than wildtype when cell lysates are treated with inorganic and organic peroxides. This is in contrast to HeLa cells, where disruption of the ESH transporter, ETT, results in an increase in protein and lipid damage [32] and ETT knockout fish, where there is an increase in levels of lipid peroxidation markers after incubation with Pb²⁺ or Cu²⁺ and an approximately four fold increase in DNA lesions in the skin [33]. The decreases in protein carbonylation, lipid peroxidation, and wildtype levels of peroxides in R119 can be explained by the dramatic upregulation of Ohr and other enzymes involved in reduction of oxidants, such as the manganese dependent catalase, KatN, and AhpC. Intriguingly, neither Ohr nor KatN are present in M. tuberculosis although there is a paralog of Ohr, OsmC (Rv2923c). Recombinant OsmC from both M. smegmatis (MSMEG_2421) and M. tuberculosis are able to reduce H2O2, CHP, and tBOH [34]. Multiple species of Ohr are present in R119, probably as a result of mycothiolation or oxidation of exposed cysteine residues resulting in the formation of sulfenic/sulfonic acids. Chi et al., 2013 [30] demonstrated that a number of proteins are mycothiolated, including the antioxidant enzymes Tpx, Gpx, and MsrA, but not Ohr, in MSH containing Corynebacterium glutamicum [30], although OhrR, the Ohr regulator, is known to be bacillithiolated in Bacillus subtilis [35]. Whether “ergothiolation” serves a similar protective role in mycobacteria remains to be seen.

5. Conclusions

Disruption of genes involved in biosynthesis of either MSH and ESH leads to up-regulation of genes involved in oxidative stress, albeit the genes that are induced are different in the two strains. The M. smegmatis MSH mutant displays protein and lipid damage despite the up-regulation of enzymes involved in protection against oxidative stress, and elevation of CoenzymeF420 levels. In contrast, the ESH mutant demonstrates less protein carbonylation and lipid peroxidation than wildtype, likely due to the up-regulation of proteins that specifically protect against oxidative stress, such as Ohr and KatN. Another key difference is result is the up-regulation of general stress proteins, GroEL and an Usps, which appears to be unphosphorylated in the MSH mutant. The lack of phosphorylation of the Usps protein in the MSH mutant suggests crosstalk between redox signaling and phosphorylation/dephosphorylation pathways in mycobacteria. Future studies are needed to determine the interplay of these two post-translational modifications in the signaling networks of mycobacteria.

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Appendix A. Transparency document

Transparency document associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.bbrep.2016.08.006.

Appendix B. Transparency document

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