Improved synthesis of the super antioxidant, ergothioneine, and its biosynthetic pathway intermediates†

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Ergothioneine and mycothiol are low molecular mass redox protective thiols present in actinomycetes, in particular mycobacteria. We report the improved chemical synthesis of ergothioneine (ESH) and biosynthetic pathway intermediates using either histidine or ESH as the starting material. The detailed mechanism of ESH biosynthesis has not yet been completely elucidated and substrates for enzymes in the pathway will provide valuable tools to aid this study. Particularly interesting is the PLP dependent β-lyase, EgtE, of mycobacteria, having the capability of cleaving the substrate, S-(β-amino-β-carboxyethyl) ergothioneine sulfoxide, to provide ESH. A synthetic route toward ESH pathway intermediates also allowed the preparation of stable isotopically labelled hercynine-d3 which was enzymatically transformed into ESH-d3. The deuterated ergothioneine biosynthetic pathway metabolites are valuable tools for future studies.

Many Gram positive bacteria, such as Mycobacterium tuberculosis, lack the redox protective molecule, glutathione, and instead produce mycothiol and ergothioneine (ESH) as their principal low molecular mass thiols.1,2 ESH is a thiohistidine betaine derivative with a thiol group at the C2 atom (ε-position) of the imidazole ring (Scheme 1). Present knowledge indicates that ESH may play a critical role in the in vivo and in vitro survival of mycobacteria.3 Recently, it was found that ESH is actively secreted into culture media by Mycobacterium smegmatis.4 A structural variant of ESH, ovothiol A, also serves as an anti-oxidant albeit in sea urchin eggs as well as in the pathogens, Leishmania major and Trypanosoma cruzi.5

Humans do not synthesize ESH, but possess an active transport system, a cation transporter (OCTN1) with high specificity for its uptake from dietary sources.6 The exact function of ESH in humans is currently in the spotlight, in particular its potent antioxidant activity.7 Recent commercial interest in ESH as a super anti-oxidant molecule has added an even greater value to improve the synthetic process development of this molecule.

In 1956, Heath et al. elucidated ESH biosynthesis in Claviceps purpurea. They demonstrated that histidine or a compound closely related to histidine might be a precursor of ESH; subsequent publications disclosed the biosynthetic assembly of ESH utilizing organisms such as Neurospora crassa and Mycobacterium smegmatis with the aid of radio isotopic labelling (14C and 35S).8–10

Melville et al. further established the participation of the S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide as an intermediate in ESH synthesis by incubation of hercynine in

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It has now been established that ESH is synthesized by the sequential action of five enzymes, encoded by the genes egtA, egtB, egtC, egtD and egtE (Scheme 1). EgtA is considered to be a γ-glutamyl cysteine ligase and catalyzes the formation of γ-glutamylcysteine. Histidine is methylated by an S-adenosylmethionine (SAM) dependent methyl transferase, EgtD, to give the trimethyl ammonium betaine, hercynine. Hercynine is then converted into S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (II) via an iron(n)-dependent oxidase (EgtB) which requires oxygen and γ-glutamylcysteine to produce γ-glutamylcysteinylhercynine (I). The exact nature of the latter transformation, in particular the sulfoxide formation, is still under investigation. Subsequently, a putative class-II glutamine amidotransidase, EgtC, mediates the hydrolysis of the N-terminus glutamic acid, providing S-γ-carboxyethyl-β-amino-γ-carboxyethyl)ergothioneine sulfoxide (II). Finally, EgtE, a pyridoxal 5-phosphate (PLP)-dependent β-lyase, gives the final product, ESH.

Recently, the research focus with regard to these mercaptohistidines has shed light on the mechanism of C–S bond formation at the δ- or ε-positions of the imidazole ring. OvoA is an iron(iii) dependent sulfoxide synthase which catalyzes the first step in ovothiol A synthesis and is a homolog of EgtB. Interestingly, the substrate specificity of EgtB vs. OvoA in achieving C–S bond formation differs significantly. OvoA is very selective towards its sulfur donor substrate and only accepts l-cysteine while it prefers histidine as the co-substrate. However, EgtB requires γ-glutamyl-l-cysteine as the sulfur donor. Furthermore, it is selective toward α-N,N,N-methylation on the histidine, i.e. hercynine as the co-substrate. Surprisingly, OvoA switches its sulfurization pattern on the histidine ring from the δ-carbon to the ε-carbon depending on the level of α-N-methylation. Thus, OvoA converts hercynine directly into S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (II) and produces a minor amount of the δ-sulfoxide (ovothiol substitution pattern) when α,N,N-dimethyl histidine is used as the co-substrate (Scheme 1).

While the enzymes EgtB and EgtC have been expressed in a functional form, EgtE is still elusive and none of the enzymes have been thoroughly studied due to the lack of readily available substrate intermediates.

Here we report the improved synthesis of the ESH and its biosynthetic precursor, S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (II). The latter sulfoxide is the substrate for the mycobacterial enzyme, EgtE. However, the absolute chirality of the sulfoxide is not known for the natural substrate or the synthetic one. Prior synthesis of intermediate (I) was reported in 1974 but was elaborate and irreproducible and resulted in a low overall yield of 8.5%. The authors reported only the position of the aromatic proton resonance and no further structural confirmation. An optical rotation, [α]D +74.4 (c = 0.5, H2O), was reported which did not reconcile with the authentic natural product [α]D +9.1 (c = 0.5, H2O). However, the m.p. of both natural and synthetic products was recorded as 188–190 °C. Nonetheless, it was claimed that synthetic S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (II) was extensively cleaved to ESH using crude cell free extracts of Neurospora crassa.

A large number of naturally occurring sulfoxides have known absolute configurations and in some instances, the differences in the biological activity of both diastereomers have been determined. S-substituted cysteine sulfoxides occur almost exclusively in the Rs-S configuration in nature. It is therefore possible that the isolated S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (II) has the Rs-S configuration. However, the latter sulfoxide’s absolute chirality and its importance are yet to be established.

Two different routes to the target compound, S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (II), were considered. In this approach, retrosynthetic cleavage of S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (II) gave the β-chloro-alanine methyl ester and ESH (Scheme 2 and S1†). Thus, S-alkylation of a protected chloromethyl alanine ester (2), derived from...
serine, provided the core structure (3). The resulting sulfide (3) was oxidised using either mCPBA or H₂O₂.

Ishikawa et al. suggested that the reaction may possibly occur via the formation of the cyclic ethylenimine carboxylic acid intermediate produced by an intramolecular S₃C₂ reaction of the β-chloroalanine (2), followed by the ring opening induced by nucleophilic attack of the sulfur atom of ESH, giving the major product N-Boc methyl ester (3a).

Sulfoxidation reaction conditions with H₂O₂ previously investigated by Ishikawa et al. led to an overoxidation to the sulfone and no analytical evidence was provided. In order to prevent overoxidation of the sulfone, mCPBA was used. Its milder nature and potential for controlled sulfoxidation compared to hydrogen peroxide is advantageous. The sulfide methyl ester (3a) was subjected to S-oxidation using one equivalent of mCPBA to afford the sulfone methyl ester (4a). The synthetic product is most likely a mixture of R₃S₃ and R₃Rₓ diastereomers. The lowest steric energy conformations with total energy 34.37 kcal mol⁻¹ of the sulfide methyl ester (3a) indicated potential face selectivity toward sulfoxidation, which could lead predominantly to the Sₙ diastereoisomer sulfoxide derivative (S1.3 & S1.4). ¹H NMR spectra of the sulfoxide methyl ester (4b) displayed evidence of diastereoselectivity (ca. 3:1 ratio) (Fig. S1.4.2 and S1.4.3f). However, only a crystal structure, supported by CD (circular dichroism) spectra, will help establish the absolute configuration of the major chiral sulfoxide and also of the natural sulfoxide (II). Deliberate oxidation of sulfide (3b), (4b) or sulfoxide (5a) to the sulfone (III) was achieved with excess oxidant.

Finally, attempted global deprotection of the Boc group and hydrolysis of the methyl ester under aqueous acidic conditions gave only the methyl ester sulfide (3b) or methyl ester sulfoxide (4b) from 3a and 4a respectively. Unsuccessful acid, base or esterase mediated ester hydrolysis obligated reconsideration of the synthetic route. Stable amino acid methyl esters have been reported before. An allyl ester derivative of β-chloroserine provided the N-Boc allyl ester sulfide (3c) after S-alkylation. Sulfoxidation followed by mild RhCl(PPh₃)₃ catalysed allyl ester cleavage and acid mediated Boc protecting group removal gave the target S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (II) in a moderate overall yield of 63%.

With the second retrosynthetic approach, cleavage of the histidine moiety of S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (3) gave cysteine and bromohercynine derivative (Scheme 2). This route has received the most attention as it provides the required sulfurization of histidine to provide the commercially important ESH. S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (5a) was synthesized in one pot using a slightly modified Erdelmeier method (Scheme 2). However, a large quantity of salt by-products had to be removed which hampered purification. Furthermore, the treatment of the S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (5a) with 3-mercaptopropionic acid at 90 °C for 18 h gave ESH. The conversion of the sulfide (5a) to the bis-benzylxoy N-Boc protected ester (5b) allowed organic extraction and removal of salts to give a clean benzyl ester (5b). Global deprotection of the N-Boc benzyl ester (5b) was achieved by hydrogenation (Pd/C) in the presence of TFA under 50 psi hydrogen pressure to give pure S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (5a). Biphasic sulfoxidation of the sulfide (5a) with mCPBA in a DCM–water mixture gave the S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (II) in a low overall yield of 36%.

The diastereomers of all S-substituted cysteine sulfoxides exhibit ¹H NMR spectra with a characteristic ABX pattern for the (S)(CH₂CH(NH₃)₂)methylene protons. ¹H NMR spectra of the S-(β-amino-β-carboxyethyl)ergothioneine sulfoxide (5a), sulfoxide (II) and sulfone (III) were consistent with the reported structures. Diagnostic ABX coupling patterns were observed for the α- and β-protons, thus giving rise to sets of doublets at about 4.0–4.5 ppm and 3.3–3.6 ppm respectively. Coupling constants Jₓₓ, Jₓₓ and Jₓₓ were approx. 10, 5 and 14 Hz respectively. ¹H NMR COSY spectra as well as ¹³C NMR aided in complete assignment of the structure.

**Improved synthesis of ESH**

Here we report the improved synthesis of the ESH via the biosynthetic precursor, S-(β-amino-β-carboxyethyl)ergothioneine sulfide. The latter sulfide and its sulfoxide (II) are substrates for the mycobacterial enzyme, EgtE, that produces ESH. Our process starts with a commercially available N-benzyl protected histidine rather than the unprotected form (key difference). We discovered that the bromination of the N-benzyl protected hercynine intermediate was achieved with N-bromosuccinimide (NBS) to give a N-debenzylated 5-bromohercynine derivative using DMF as solvent in 90% yield. The advantage of the latter intermediate is that subsequent process steps are almost quantitative, relatively simple, all at room temperature, shortened and allow an overall synthesis yield of 70%. Thus, our process is at least 2 times better in overall yield than any prior patented or published process. The high yields of the intermediate products also allow viable isotopic labeling steps to be performed. Isotopes are usually very expensive and are advantageous by high yield conversions. The final step involves a biomimetic pyridoxal phosphate (PLP) mediated cleavage of the sulfide or sulfoxide substrates with crude enzymatic extracts of *M. smegmatis* to give ESH (Scheme 3).

Previously, the biosynthesis pathway of ESH was elucidated utilising radiolabeled intermediates. Advances in synthetic procedures have now made it possible to synthesize the same intermediates incorporating stable isotopes. These intermediates are valuable internal standards in the quantitation of pathway metabolites during external stimuli or drug treatment. Here we also report the synthesis of ESH-d₁₀ (10) and demonstrate its biosynthesis from deuterated hercynine.

Hercynine-d₁₀ (7) was synthesized in a two-step reaction starting with the commercially available L-histidine (Scheme 2). The first step involved reductive amination using aqueous formaldehyde and sodium triacetoxyborohydride to give N,N-dimethyl histidine (6). The second step involved the quaternarization of the crude N,N-dimethyl histidine (6) using
methyl-d₃ iodide under basic conditions to give the hercynine-

d₃ (7). Characteristic HRMS peaks indicating M + 3 for hercynine-
d₃ (7) were obtained (S3.2.1†).

ESH-d₃ (10) was synthesized in two sequential reaction
steps starting with the S-tert-butyl protected 2-mercaptop-
histidine (9), derived from mercaptohistidine (8). Selective
N-methylation with methyl-d₃ iodide, followed by S-tert-butyl
de-protection using 2-mercaptoacetic acid (tert-butyl scavenger)
in HCl gave ESH-d₃ (10). Characteristic HRMS peaks indicating M + 3 for the ESH-d₃ (10) were obtained (S3.2.2†).

We compared the enzymatic and non-enzymatic PLP
mediated synthesis of ESH from the synthetic sulfoxide (II). To
this end, crude M. smegma-
tis cell free extracts were isolated from cultures grown and harvested at the late exponential
phase, characterised by high enzymatic activity.10

The crude enzymatic transformation of the ESH bio-
synthetic pathway precursors, including sulfide and sulfoxide
variants, was evaluated by the concomitant production of ESH
in excess of basal levels as determined by LCMS. ESH precu-
sor metabolites hercynine-d₃ (7), hercynyl cysteine methyl ester
sulfoxide (4b), (β-amino-β-carboxyethyl)ergothioneine sulfoxide
(15) and (β-amino-β-carboxyethyl)ergothioneine sulfoxide (II)
were incubated with the crude cell free extract at 37°C, pH = 7.4 for 1 day, and analyzed by LCMS.

The control reaction containing only the crude M. smegma-
tis cell free extract was also treated under the same conditions
as the four metabolites. The concentration of ESH thus
obtained was 5.70 (±0.30) ng ml⁻¹, which equate to that of
endoogenous ESH. This concentration was above the limit of
detection (0.78 ng ml⁻¹), thus any increase in the concen-
tration of ESH in the experiment above 1 ng ml⁻¹ is considered
significant enough to be ascribed to basal levels or bio-
transformation of the respective substrates by the crude
endoogenous enzymes of the ESH pathway.

When hercynine-d₃ (7) was used as the substrate, as
expected, no change in baseline ESH was observed. However,
as expected, the LCMS analysis of the mixture revealed the pro-
duction of ESH-d₃ (10). The HRMS (ESI⁺) displayed a peak at
m/z 233.1161 corresponding to [M + d₃]⁺ (S4.2.1†). Peak values
at m/z 230.0958 [M⁺] or at m/z 231.0980 [M + H⁺] belonging to
the natural ESH were not present in the chromatogram. This
clearly shows that the crude M. smegmatis enzymes (dialyzed
transform) the hercynine-d₃ (7) into ESH-d₃ (10). This
proved that the isolated crude enzyme extract was fully func-
tional in the reconstitution of ESH synthesis.

(β-Amino-β-carboxyethyl)ergothioneine sulfoxide (II)
bioc- synthetically produced the highest concentration of ESH
(22.6 ng ml⁻¹) (Fig. 1). The (β-amino-β-carboxyethyl)ergothio-
ene sulfide (15) appeared to be almost as good a substrate as
the (β-amino-β-carboxyethyl)ergothioneine sulfoxide (II)
(19.2 ng ml⁻¹ ESH). It is well known that PLP-dependent trans-
formations can also undergo enzyme free conversion albeit
with a much slower rate and specificity.22 Thus, the non-enzym-
atic treatment of the sulfide (15) with 50 mM PLP at 37°C
resulted in an efficient formation of ESH (96.3 ng ml⁻¹)
(Fig. S4.5.1†). However, under the same conditions, the sulfox-
ide (II) produced no ESH at all (Fig. S4.5.2†).

The pKₐ value of the amino acid ε-hydrogen usually is in
the range 20–30 and many enzymes effectively increase the
ε-hydrogen acidity with the aid of a coenzyme, PLP. PLP-dependent enzymes exist in their native state as an internal
aldehyde (Schiff base) (S-5.1–5.3‡) with the ε-amino group of a
lysin residue present in the catalytic site. The amino acid sub-
strate displaces the lysine from the internal aldehyde to form
a new aldehyde, termed external aldehyde. The formation of
such an external aldehyde can reduce the pKₐ value of the
ε-proton of the substrate from 30 to as low as 6.22

PLP-dependent enzymes have been differentiated on the
basis of their tertiary structures, more specifically according to
fold types I–V. EgtE is classified as a fold type V which is found mainly in enzymes that catalyze β-replacement and β-elimination reactions. In the absence of a crystal structure of EgtE, structure homology analysis amongst mycobacteria is not possible. However, a putative β-lyase from *E. tasmaniensis* appeared to catalyze this step but has a sequence similarity to EgtE of *M. Smeg* and *M. tb* of only 14%.12 Thus, the proposed mechanism of this C-S β-lyase utilizing (β-amino-β-carboxyethyl)ergothioneine sulfide (15) as a substrate was based on that of the well characterized cystalysin from *Trepnoma denticola* which shares 31% sequence identity to the β-lyase from *E. tasmaniensis*.23 It may well be that actinomycetes have similar β-lyases at work.24

We also observed a rather facile non-enzymatic (PLP only) β-elimination with (β-amino-β-carboxyethyl)ergothioneine sulfide (15). However, the conversion of the natural substrate, (β-amino-β-carboxyethyl)ergothioneine sulfoxide (II), to ESH is not that straightforward. Elimination of an unstable (β-amino-β-carboxyethyl)ergothioneine sulfinate (IV) is envisaged, whereby self-condensation leads to the thiosulfinate, which in turn decomposes to ESH and an equivalent amount of relatively stable (β-amino-β-carboxyethyl)ergothioneine sulfinic acid (V). The latter sulfinic acid can subsequently be reduced to the thiol, ESH, with the aid of excess mercaptoethanol, but this may be difficult under the current experimental conditions.25 Note that the (β-amino-β-carboxyethyl)ergothioneine sulfinate (II) did not produce ESH (in the absence of mercaptoethanol).

Efforts are underway to purify and crystallize EgtE, which would allow a better understanding of enzyme–substrate binding, specificity and the potential for inhibitor design. An EgtD deletion mutant of *M. Smegmatis* and a mycothiol-deficient mutant did not affect their susceptibility to antibiotics.4 However, the ESH/mycothiol-deficient double mutant was significantly more sensitive to peroxide than either of the single mutants lacking either ESH or mycothiol, suggesting that both thiols play a role in protecting *M. Smegmatis* against oxidative stress. Thus, an inhibitor of ESH synthesis will be valuable in drug susceptibility studies of mycobacteria.

**Competing interest**

The authors declare no competing financial interest.

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**References**