ASSAY FOR IDENTIFYING INHIBITORS OF MYCOBACTERIUM ANTI-OXIDANT DEFENSE SYSTEM

Abstract: The invention relates to an enzyme-based assay for detecting inhibitors of the anti-oxidant defense system in Mycobacterium involving a thioredoxin reductase or an alkylhydroperoxidase, a method for manufacturing a pharmaceutically active compound for treating and/or preventing a Mycobacterium associated disease and uses thereof.
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ASSAY FOR IDENTIFYING INHIBITORS OF MYCOBACTERIUM
ANTI-OXIDANT DEFENSE SYSTEM

Description

Field of the invention

The present invention pertains generally to an enzyme-based assay for detecting inhibitors of the anti-oxidant defense system in Mycobacterium. In a first aspect, the enzyme-based assay comprises a reductant, a thioredoxin reductase or a peroxidase, a donor substrate capable of receiving a hydrogen atom from said thioredoxin reductase or said peroxidase, and a hydroperoxide. The enzyme-based assay is capable for determining the consumption of the reductant.

Furthermore, the invention relates to a method for manufacturing a pharmaceutically active compound for the treatment and/or the prevention of a Mycobacterium associated disease. In a first aspect, the method comprises the steps of providing an enzyme-based assay for detecting inhibitors of the anti-oxidant defence system in Mycobacterium, providing a test sample comprising a potentially active compound, contacting the enzyme-based assay with the test sample, determining the consumption of a reductant and recovering the active compound.

Additionally, the invention refers to a pharmaceutically active compound, which is identified by the method for manufacturing said pharmaceutically active compound. Said compound is used for the preparation of a medicament for treating and/or preventing a Mycobacterium associated disease.
Several documents are cited throughout the text of this specification. Each of the documents cited herein (including any manufacturer’s specifications, instructions etc.) are hereby incorporated by reference. However, there is no admission that any document cited is indeed prior art to the present invention.

**Background of the invention**

*Mycobacteria* cause two of the most important diseases in history, tuberculosis and leprosy. Even though the antibiotic streptomycin, which possesses a broad spectrum of activity, is usually active against *Mycobacterium tuberculosis*, many clinical isolates have been shown to be resistant. Further, resistance of *Mycobacterium tuberculosis* to the tuberculostatic isoniazid (INH) is detected in 4-8% of clinical isolates in the United States and is much higher in other populations (Pablos-Mendez et al., (1998), *N. Engl. J. Med.* 338, 1641-1649). It is assumed that INH resistance is at least in part due to a loss-of-function mutation of a key enzyme in the metabolic pathway.

Bacteria, as living organisms need energy to survive. This energy is produced by a process known as oxidative phosphorylation, in form of the chemical entity adenosine triphosphate (ATP). A major role in the production of ATP play two pyrimidine nucleotides, namely nicotinamide adenine dinucleotide (NAD) and nicotinamide adenine dinucleotide phosphate (NADP). These compounds mediate most often biological oxidations and
reductions, i.e., they serve as acceptors for hydrogen atoms released by dehydrogenation reactions, and as donors of hydrogen atoms required for hydrogenation reactions. Both of these pyrimidine nucleotides can readily undergo reversible oxidation and reduction.

Bacteria do not only have to produce energy in order to survive, but they also have to deal with oxygen. All bacteria contain certain enzymes, which react with oxygen, e.g., superoxide dismutase, catalase and peroxidase. These enzymes protect the cell from the toxic consequences of oxygen metabolism, as they eliminate oxygen radicals and hydrogen peroxide ($H_2O_2$).

Organisms living under aerobic conditions have developed various anti-oxidative mechanisms to protect themselves from damage by reactive oxygen species. A family of anti-oxidative proteins, designated as peroxiredoxin, has been identified. They share a common reactive cysteine residue in the N-terminal region, and are capable of serving as a peroxidase and involve thioredoxin and/or glutathione as the electron donor.

Peroxiredoxins are low efficiency peroxidases using thiols as reductants. They appear to be fairly promiscuous with respect to the hydroperoxide substrate; the specificity for the donor substrate varies considerably between the subfamilies, comprising GSH, thioredoxin, tryparedoxin and the analogous CXXC motifs in bacterial AhpF proteins. Peroxiredoxins are responsible for anti-oxidant defense in bacteria (AhpC), yeast (thioredoxin peroxidase) and trypanosomatids (tryparedoxin
peroxidase). They are considered to determine virulence of mycobacteria.

Thioredoxins are small molecular weight disulfide oxidoreductases specialized in the reduction of disulfide bonds on other proteins. Generally, the enzymes which are selectively and reversibly reduced by these proteins oscillate between an oxidized and inactive conformation and a reduced and active conformation. Thioredoxin constitutes the archetype of a family of protein disulfide oxidoreductase which comprises glutaredoxin and protein disulfide isomerase. Thioredoxin and glutaredoxin serve many roles in the cell, including the redox regulation of target enzymes and transcription factors. They can also serve as hydrogen donors to peroxiredoxins, the function of which is to get rid of hydroperoxide in the cell.

Isoniazid (INH) resistance of Mycobacterium tuberculosis is supposed to be due to a loss-of-function mutation of catalase (katG) (Sherman et al., (1996), Science 272, 1641-1643) an enzyme, which is required for the activation of INH (Zhang et al., (1992), Nature 358, 591-593), but appears equally important for removal of H2O2. Catalase-negative clinical isolates constitutively overexpress AhpC, which is a two-cysteine-peroxiredoxin-type peroxidase, whereas catalase containing strains (katG+ strains) generated in vitro, which do not overexpress AhpC, proved to be virulent. This revealed that the catalase deficiency had to be compensated by another peroxidase, such as AhpC, to assure protection of the pathogen against the oxidant attack of the host's phagocytes (Sherman et al., (1999), Biofactors 10, 211-217). However, it is unclear in the prior art how AhpC is supplied with reduction

Several publications deal with agents which are useful in the treatment of mycobacterial infections, like tuberculosis. Among them, for example WO 03/004479 A1 discloses therapeutic agents, in particular certain 4-aryl-cyclohexa-2,5-dienone compounds (4-aryl quinols), which are supposed to be effective in the treatment of proliferative conditions, cancer, mycobacterial infections (e.g. tuberculosis), and/or conditions mediated by thioredoxin/thioredoxin reductase. The application shows that 4-aryl quinols are inhibitors of microbial growth and, in addition, inhibit the human thioredoxin/thioredoxin reductase system. However, this application does not provide for an *in vitro* system based on components derived from the mycobacterial peroxidase system.

The resistance of *Mycobacterium tuberculosis* to isoniazid is commonly linked to inactivation of the catalase-peroxide KatG that converts isoniazid to its biological active form. Loss of KatG is associated with elevated expression of the alkyl hydroperoxidases AhpC and AhpD. AhpD has no sequence identity with AhpC or other proteins but has alkyl hydroperoxidase activity and possibly additional physiological activities. The alkyl hydroxyperoxidase activity, in the absence of KatG provides an important antioxidant defense. Therefore, Nunn et al., (2002), *J. Biol. Chem.* 277, 20033-20040 determined the *M.*
tuberculosis AhpD structure to a resolution of 1.9 Å. The protein was found to be a trimer in a symmetrical cloverleaf arrangement. Each subunit exhibits a new all-helical protein fold in which the two catalytic sulfhydryl groups, Cys-130 and Cys-133, are located near a central cavity in the trimer. Based on the crystal structure, the authors suggest a catalytic mechanism for AhpD involving a proton relay in which the Glu-118 carboxylate group, via His-137 and a water molecule, deprotonates the catalytic residue Cys-133.

In order to test this suggested mechanism, Koshkin et al., (2003), J. Biol. Chem. 278, 29502-29508 expressed several mutants of AhpD, determined the crystal structures of the H137F and H132Q mutants, estimated the pKa values of the cysteine residues, and defined the kinetic properties of the mutant proteins. Their results supported the proposed catalytic mechanism for AhpD.

Zhang et al., (1999), Archives of Biochemistry and Biophysics 363, 19-26 expressed the thioredoxin (Trx) and thioredoxin reductase (TR) of Mycobacterium tuberculosis in E. coli and demonstrated the reduction of peroxides and dinitrobenzenes. The reduction of H2O2 required both Trx and TR and was more efficient under anaerobic than aerobic conditions. In contrast, cumene hydroperoxide is reduced to cumyl alcohol and acetophenone in a process that requires NADPH and TR but not Trx.

WO 96/19578 A2 discloses the identification and characterization of the ahpCF operon that determines mycobacterial resistance to the antibiotic isoniazid and its analogues. The operon encodes the polypeptides AhpC and AhpF, which are sup-
posed to combine to form an active alkyl hydroperoxide reductase enzyme that may either be a direct target of INH or act to confer INH resistance. The sequence of a mutant ahpCF operon is provided, showing that INH resistance can be conferred by a mutation in the promoter region. Also provided are polynucleotides and polypeptides that are useful in diagnosis and treatment.

None of the above cited prior art documents discloses an efficient in vitro test system composed only of component which are indeed present in Mycobacterium. The test system aims to block the system in Mycobacterium rather than to block any artificial peroxidase or thioredoxin as disclosed and suggested in the prior art. The inventors of the present invention provide for the first time a test system in which only components of the mycobacterial anti-oxidant defense system co-operate in situ without the help of any artificial protein.

As stated above, many clinical isolates of Mycobacterium are resistant to commercially available antibiotics. This is not only a problem with respect to an adequate treatment of infected patients, who bear the risk of developing a chronic infection, but it is also a serious problem for test systems, which are currently used to identify strains and clinical isolates in samples. In order to develop new highly efficient antibiotic compounds, there is a need for an assay system, which is capable of identifying such active compounds.
Summary of the invention

The present invention relates to a novel enzyme-based assay for detecting inhibitors of the anti-oxidant defense system in Mycobacterium. Furthermore, the present invention relates to a method for manufacturing a pharmaceutically active compound for treating and/or preventing a Mycobacterium associated disease. The enzyme-based assay and the method for manufacturing a pharmaceutically active compound are useful for the identification of compounds active in the treatment and/or prevention of Mycobacterium associated diseases. With the help of such pharmaceutically active compounds, diseases like tuberculosis caused by antibiotic resistant strains, can be efficiently treated and/or prevented. Therefore, the invention additionally relates to the use of a pharmaceutically active compound identified by a method for manufacturing pharmaceutically active compounds for the preparation of a medicament for treating and/or preventing a Mycobacterium associated disease.

Therefore, the present invention provides for the first time an enzyme-based assay in the form of a test system in which only components of the mycobacterial anti-oxidant defense system co-operate in situ without the help of any artificial protein (not derived or naturally occurring in Mycobacterium).

Brief description of the drawings

The disclosure of this invention may best be understood in conjunction with the accompanying drawings, in which
Figure 1 shows oxidative activation of Mycobacterium tuberculosis AhpC (MtAhpC) and Mycobacterium tuberculosis AhpD (MtAhpD) by t-butyl hydroperoxide (t-booH). MtAhpC activity with MtAhpD as cosubstrate was tested in the conventional test system.

Figure 2 shows a steady-state-kinetic analysis of Mycobacterium tuberculosis peroxiredoxin TPx (MtTPx) with Mycobacterium tuberculosis thioredoxin B (MtTrxB) or Mycobacterium tuberculosis thioredoxin C (MtTrxC) as cosubstrates.

**Detailed description of the invention**

In view of the need of therapeutic means for the prevention and treatment of diseases related to mycobacteria, the technical problem underlying the invention is to provide means and methods for the identification of pharmaceutically active compounds, which act as inhibitors of the anti-oxidant defense system in Mycobacterium.

The solution to the technical problem is achieved by providing following embodiments of the present invention.

Accordingly, in a first aspect the invention relates to an enzyme-based assay for detecting inhibitors of the anti-oxidant defense system in Mycobacterium comprising

(i) a reductant;
(ii) a thioredoxin reductase or a peroxidase;
(iii) a donor substrate capable of receiving a hydro
gen atom from (ii); and
(iv) a hydroperoxide,
for determining the consumption of the reductant.

In a preferred embodiment of the present invention, the thioredoxin reductase and/or the peroxidase and/or the donor substrate is an enzyme derived from Mycobacterium. A particularly preferred bacterium is Mycobacterium tuberculosis.

In a particularly preferred embodiment, all components of the enzyme-based assay are simultaneously derived from Mycobacterium. Again, a particularly preferred bacterium is Mycobacterium tuberculosis.

In another embodiment, the enzyme is expressed in heterologous host cells. Preferred host cells are cells of bacteria, yeasts, insect cells or mammalian cells. Particularly preferred are cells of Escherichia coli, Saccharomyces cerevisiae and Spodoptera frugiperda.

The reductant, which is used in the enzyme-based assay is preferably NADH, NADPH and/or FADH.

In another preferred embodiment of the present invention, the peroxidase, which is used in the enzyme-based assay, is a peroxiredoxin. Preferred are AhpC and/or TPx of Mycobacterium, and enzymes of Mycobacterium tuberculosis is most preferred.

The donor substrate, which is used in the enzyme-based assay is preferably glutathione or thioredoxin. Preferred are thio-
redoxin B (TrxB) and/or C (TrxC) of *Mycobacterium*, and enzymes of *Mycobacterium tuberculosis* are most preferred.

The hydroperoxide, which is used in the enzyme-based assay, can be hydrogen peroxide, t-butyl hydroperoxide, cumene hydroperoxide, linoleic acid hydroperoxide, phosphatidyl choline hydroperoxide or any oxidant comprising a hydroperoxy group.

In another aspect, the present invention relates to a method for manufacturing a pharmaceutically active compound for treating and/or preventing a *Mycobacterium* associated disease, comprising the steps of

(a) providing an enzyme-based assay comprising a reductant, a thioredoxin reductase or a peroxidase, a donor substrate capable of receiving a hydrogen atom from the thioredoxin reductase or the peroxidase, and a hydroperoxide;

(b) providing a test sample comprising a potentially active compound;

(c) contacting (a) and (b);

(d) determining consumption of a reductant;

(e) recovering the active compound.

In a preferred embodiment, the reductant used in the method for manufacturing a pharmaceutically active compound is NADH, NADPH and/or FADH.

In another embodiment, the active compound is combined with suitable pharmaceutically acceptable fillers, excipients and/or additives. Preferred are physiological saline solution,
demineralised water, stabilizers, proteinase inhibitors and gel formulations.

In a third aspect, the present invention relates to the use of the pharmaceutically active compound identified by the method for manufacturing a pharmaceutically active compound, for the preparation of a medicament for treating and/or preventing a disease, which is associated with Mycobacterium.

In a preferred embodiment, the disease is tuberculosis, which is associated with Mycobacterium tuberculosis.

**Best method to carry out the invention**

Thioredoxin C that, by analogy to many eukaryotic systems, was suspected to reduce peroxiredoxins. However, it was reported to be inactive as potential alternative reductant of mycobacterial AhpC (Hillas et al., (2000), J. Biol. Chem. 275, 18801-18809), as was mycothiol, the major low molecular weight thiol of mycobacteria (Vergauwen et al., (2001), Arch. Physiol. Biochem. 113, B22).

Even though thioredoxin C was convincingly demonstrated to be inactive in reducing AhpC, the inventors screened the genome of Mycobacterium tuberculosis (Cole et al., (1998), Nature 393, 537-544) for homologues of AhpF, thioredoxin-related proteins and peroxiredoxins. Subsequently, they expressed the genes heterologously in Escherichia coli or Mycobacterium smegmatis, and were surprisingly able to reconstitute peroxidase systems from the isolated gene products in vitro. Fur-
ther, the inventors checked the efficiencies of the heterolo-
gously expressed proteins by steady-state kinetics.

The inventors demonstrated that mycothiol reductase of
(MtMycR), which is a homologue of AhpF, does not reduce
MtAhpC. Neither did mycothiol if continuously regenerated by
NAD(P)H via MtMycR.

Further, the inventors demonstrated that MtAhpD is reduced by
liponamide and can in turn reduce MtAhpC. Liponamide, kept re-
duced by NADH and bovine liponamide reductase, plus MtAhpD and
MtAhpC constituted a peroxidase system. The time progression
curves, however, were unusual in starting with a lag phase
when the fully reduced system was started with t-bOOH as
MtAhpC substrate (Figure 1). Reversal inhibition by the ini-
tial high hydroperoxide concentration does not account for the
phenomenon, since it is identically observed from 20 μM up to
73 μM initial t-bOOH.

Surprisingly, the reaction started at maximum rate if MtAhpC
and MtAhpD were preincubated with t-bOOH for 2 min, indicating
a facilitated interaction of the reaction partners. The x-ray
structure of AhpD shows that the reactive CXXC motifs of AhpD
are hidden in the interior of the trimeric protein (Bryk et al.,
(2002), Science 295, 1073-1077; Nunn et al., (2002), J.
Biol. Chem. 277, 20033-20040), and MtAhpC tends to form
decameric rings in dependence of its redox status (Chauhan and
Biochemistry 41, 5493-5504). Redox-dependent conformational
changes might therefore determine the accessibility of the
MtAhpD reaction centers for MtAhpC. Steady-state-analysis of
the apparently regular parts of the turnover curves yielded a net forward rate constant of MtAhpC for the reaction with t-bOOH, k'₁, of 1 \times 10^4 \text{ M}^{-1}\text{s}^{-1}, which marks the lower edge of corresponding peroxiredoxin constants (Hofmann et al., (2002), supra; Wood et al., (2003), supra), and a k'₂ for the reduction by MtAhpD of 2.7 \times 10^6 \text{ M}^{-1}\text{s}^{-1} which, like the K_M of 0.23 \text{ µM}, clearly demonstrates specific reactivity.

Additionally, the inventors were surprisingly successful in reconstituting mycobacterial peroxidase systems in vitro by combining AhpC with thioredoxin reductase (MtTR) and various CXXC-containing proteins. MtTR did not reduce MtAhpD, MtTrxA and the glutaredoxin related Rv2466c gene product but only MtTrxB and MtTrxC. Surprisingly, MtTrxC efficiently reduced MtAhpC. The kinetic analysis of MtAhpC in the system reconstituted by homologous TR and TrxC yielded a k'₁ of 2.3 \times 10^4 \text{ M}^{-1}\text{s}^{-1} that is nearly identical to that obtained in the AhpD system. Both the k'₂ and the K_M characterizing the reactivity and/or affinity to TrxC compare unfavourably with the constants for AhpD by a factor of 10 and 20, respectively, as does the k_{cat} (Table 1), yet the relative cellular abundances of the potential AhpC reductants (see below) qualify MtTrxC as the relevant one.

MtTrxC also proved to reduce another mycobacterial peroxiredoxin (Rv1932) that by sequence homology, had been classified as a “thiol peroxidase” (TPx) but had never been investigated in respect to activity and specificity. MtTPx also accepted MtTrxB although with slightly lower affinity and reactivity (Table 1). MtTPx belongs to a subfamily of two-cysteine-peroxiredoxins which, in contrast to AhpC and other two-
cysteine-peroxiredoxins, are considered to be functionally monomeric. Their mechanism of action is believed to involve oxidation of an N-proximal cysteine residue, here C60, to a sulfenic acid derivative by the hydroperoxide substrate followed by formation of an intramolecular disulfide bridge with a distal cysteine, here C94, while in the oligomeric two-cys peroxiredoxins the corresponding reaction center is built up between the subunits (Hofmann et al., (2002), supra; Wood et al., (2003), supra; Baker and Poole, (2003), J. Biol. Chem. 278, 9203-9211).

Steady-state kinetics of MtTPx with t-butyl hydroperoxide and MtTrxB as substrates display a ping-pong pattern, as is typically observed with peroxidases (Figure 2A, B). With MtTrxC as substrate, however, the slopes in reciprocal primary plots (Figure 2C) surprisingly appear not strictly parallel and show marked deviations from linearity at high peroxide concentrations that are particularly pronounced at low cosubstrate levels. This kinetic anomaly is also seen with AhpC. This phenomenon is interpreted as being indicative of negative cooperativity between subunits. Gel permeation of native MtTPx yielded an average size of 36 kDa that underscores its homodimeric nature. The kinetic anomalies of MtTPx could thus be due to an allosteric phenomena.

Interestingly, the peroxidatic efficiency of MtTPx is about one order of magnitude higher than that of MtAhpC. With MtTrxB as cosubstrate a $k'_1$ value $3.3 \times 10^5 \text{ M}^{-1}\text{s}^{-1}$ was measured that appears more reliable than the corresponding value obtained with MtTrxC of $0.9 \times 10^5 \text{ M}^{-1}\text{s}^{-1}$. In a ping-pong mechanism, the values should be identical but the latter is likely un-
derestimated due to the deviations from linearity of the reciprocal plots (Figure 2C). The $k'_2$ values of MtTPx for both thioredoxins are similar and slightly higher than that of MtAhpC for MtTrxC but lower than that of the MtAhpC/MtAhpD couple. The highest $V_{\text{max}}$ is obtained for the MtTPx/MtTrxC system (Table 1).

The kinetic constants compiled in table 1 and the general rate equation for two-substrate ping-pong mechanisms

$$[E_0]/v = \Phi_0 + \Phi_1/[\text{ROOH}] + \Phi_2/[\text{reductant}]$$

allow to estimate turnover rates by the different systems if, as a first approximation, the peculiarities of peroxiredoxin kinetics at extreme substrate concentrations are ignored. However, the concentrations of the enzymes appear to vary between strains and substrate concentrations can only be roughly estimated. At low peroxide tone and high supply of reductants, as is commonly assumed for unstressed situations, the turnover depends on the level of reduced peroxidase that should approximately equal $[E_0]$, on $k'_1$ and on the peroxide concentration.

$$v = k'_1 [E_{\text{red}}] [\text{ROOH}]$$

AhpC is hardly detectable in most of the $\text{katG}^+$ strains of $\text{Mycobacterium tuberculosis}$ and $\text{Mycobacterium marinum}$ while it is easily found in $\text{Mycobacterium smegmatis}$ and $\text{Mycobacterium bovis}$ BCG. In contrast, MtTPx is consistently seen in proteomes of $\text{Mycobacterium tuberculosis}$. Its concentration may be estimated much higher than the concentration of AhpC. Being the
dominant peroxiredoxin and having a higher $k'_1$ than AhpC, MtTPx must definitely be rated as the more relevant enzyme under physiological conditions. Under $H_2O_2$ stress, the reductant capacity may become limiting according to

$$v = k'_2 [E_{ox}] [\text{reductant}]$$

The concentration of the reductants is linked in a complex manner to the NADH and NADPH pools, depends on the speed of regenerating enzymes and the pool size of the proximal reductants, only the latter being easily estimated. AhpD can be identified in *Mycobacterium tuberculosis* proteomes as a minor spot corresponding to a low molarity level. MtTrxB is characterized here as a specific substrate of MtTPx, but MtTrxC is always seen as a major protein in all *Mycobacterium tuberculosis* strains investigated. MtTR is a medium spot in all analyzed mycobacterial proteomes (Jungblut et al., (1999), *Mol. Microbiol.* 33, 1103-1117; Mollenkopf et al., (1999), *Electrophoresis* 20, 2172-2180; Nagai et al., (1991), *Infect. Immun.* 59, 372-382; Sonnenberg and Belisle, (1997), *Infect. Immun.* 65, 4515-4524; Weldingh et al., (1998), *Infect. Immun.* 66, 3492-3500; Rosenkrands et al., (2000), *Electrophoresis* 21, 935-948; Rosenkrands et al., (2000), *Electrophoresis* 21, 3740-3756. Thus, the MtTR/TrxC-mediated detoxification systems, because of their high reduction capacities, are particularly relevant under conditions of oxidative stress.

Taking into account both rate constants and capacities, NADPH-driven $H_2O_2$ reduction by the MtTR/MtTrxC/MtAhpC and the MtTR/MtTrxC/MtTPx system should be at least as efficient as the NADH-driven, AhpD/AhpC-mediated one in coping with $H_2O_2$. 
challenge into $\text{katG}^-$ strains. Both peroxiredoxin-type peroxidases may, however, meet special antioxidant requirements also in $\text{katG}^+$ strains. Like other peroxiredoxins, the mycobacterial enzymes act on a broad spectrum of hydroperoxides, $\text{MtTPx}$ even on complex lipid hydroperoxides (Table 2). $\text{MtAhpC}$ and $\text{MtTPx}$ therefore complement catalase in antioxidant defense of $\text{Mycobacterium tuberculosis}$ as do broad spectrum selenoperoxidases, such as GPx-4, in the host (Flohé and Brigelius-Flohé, (2001), in: Selenium. Its Molecular Biology and Role in Human Health, D. L. Hatfield, Ed. (Kluwer Academic Publishers, Boston/Dordrecht/London), and thereby contribute in general to the pathogen’s resistance to the oxidative stress exerted by the innate immune response.

**Examples**

The invention will now be described in detail with respect to showing how certain specific representatives embodiments thereof can be implemented. However, all examples are intended to be illustrative only. In particular, the invention is not intended to be limited to method, materials, conditions, process parameters and the like specifically recited herein.

**Example 1**

**Preparation of constructs**

$\text{MtMycR preparation}$
The MycR open reading frame (ORF) was amplified from *Mycobacterium tuberculosis* H37Rv genomic DNA by PCR with engineered 5' *PstI* and 3' *HindIII* sites and a C-terminal 6xHis tag (*MtMycR* Forward: 5'-NNN NNN CTG CAG ATG GAA ACG TAC GAC ATC GCG ATC-3'; *MtMycRH6* Reverse: 5'-NNN NNN AAG CTT TCA GTG ATG GTG ATG ATG ACG CAG GCC AAG CAG CGC GTT-3'). The PCR product was cloned into the mycobacterial expression vector pMV261. Protein expression was induced in *Mycobacterium smegmatis* by heat shock (45°C). The soluble protein was purified by nickel chelate chromatography (Ni-NTA Superflow; Qiagen, Hilden, Germany).

**MycSH preparation**

*Mycobacterium smegmatis* cells were disrupted by boiling them for 10 min at 95°C in 10 mM Tris-HCl pH 8. MycSH was isolated from the crude *Mycobacterium smegmatis* soluble extract essentially as described by Unson et al., (1998), *J. Immunol. Methods* 214, 29-39.

**MtAhpC preparation**

The AhpC ORF was amplified by PCR using a forward primer that contained an *NdeI* site and overlapped the 5' end of the coding sequence (*MtAhpC* Forward: 5'-G GCG CGC CAT ATG CCA CTG CTA ACC ATT GGC GA-3') and a reverse primer which contained a *BamHI* site, the sequence for a 6xHis tag and overlapped the 3' end of the coding sequence (*MtAhpCH6* Reverse: 5'-CCG GCG GGA TCC TTA GTG GTG GTG GTG GTG GCC CGA AGC CTT GAG GAG TT-3').
The PCR product was cloned into the procaryotic expression vector pET22b(+) (Novagen, Madison, USA) digested with corresponding sets of enzymes. Expression was induced in *Escherichia coli* Tuner (DE3) (Novagen, Madison, USA) with 0.5 mM IPTG at 30°C. The soluble protein was purified by nickel chelate chromatography.

**MtAhpD preparation**

The AhpD ORF was amplified by PCR using primers with engineered 5' NdeI and 3' HindIII sites (MtAhpD Forward: 5'-CC CCC CCC CAT ATG AGT ATA GAA AAG CTC AAG GCC GCG-3'; MtAhpD Reverse: 5'-CC CCC CCC AAG CTT TTA GCT TGG GCT TAG TGC CTC GAT-3') and cloned into pET22b(+). Expression was induced in *E. coli* Tuner (DE3) with 0.5 mM IPTG at 30°C. MtAhpD was purified to homogeneity by anion exchange chromatography (UNOsphere Q; BioRad, Munich, Germany) and hydrophobic interaction chromatography (Phenyl Sepharose 6; Amersham Biosciences, Freiburg, Germany).

**MtTR preparation**

The vector pTrcHis A (Invitrogen, Karlsruhe, Germany) containing the TR ORF fused to an N-terminal 6xHis tag was kindly provided by T.H.M. Ottenhoff (Leiden University Medical Centre, Leiden, The Netherlands). Protein expression and purification was done as described for MtAhpC.
MtTrxA, MtTrxB and MtRv2466c preparation

The TrxA, TrxB and Rv2466c ORFs were amplified by PCR using forward primers that contained an NdeI site and overlapped the 5' end of the coding sequence (MtTrxA Forward: 5'-CCC CCC CAT ATG GTG ACC ACT CGA GAC CTC ACG-3' ; MtTrxB Forward: 5'-CCC CCC CAT ATG GTG ACT ACC CGA GAC CTC ACT GCC-3' ; MtRv2466c Forward: 5'-CCC CCC CAT ATG CTC GAG AAG GCC CCC CAG-3') and reverse primers which contained an EcoRI site, the sequence for a 6xHis tag and overlapped the 3' end of the coding sequence (MtTrxAH6 Reverse: 5'-CCC CCC GAA TTC TCA GTG ATG ATG ATG ATG ATG GGA TGA AGT CTT TGT TCC AGG GCC-3' ; MtTrxBH6 Reverse: 5'-CCC CCC GAA TTC TCA GTG ATG ATG ATG ATG ATG GGC TTG TTG GGC TCG CCC GTT-3' ; MtRv2466cH6 Reverse: 5'-CCC CCC GAA TTC CTA GTG GTG GTG GTG GTG GTC GAA CTG AGG CGG CTC GGT-3'). The PCR products were cloned into pET22b(+) digested with corresponding set of enzymes. Expression and purification was done as described for MtAhpC.

MtTrxC and MtTPx preparation

The TrxC and TPx ORFs were amplified by PCR using forward primers that contained an NdeI site, the sequence for a 6xHis tag and overlapped the 5' end of the coding sequence (MtH6TrxC Forward: 5'-CC CCC CCC CAT ATG CAT CAC CAT CAC CAT CAC ACC GAT TCC GAG AAG TCC GCC-3' ; MtH6TPx Forward: 5'-CCC CCC CAT ATG CAT CAC CAT CAC GCA CAG ATA ACC CTG CGA GGA-3') and reverse primers which contained a HindIII site and overlapped
the 3' end of the coding sequence (MtTrxC Reverse: 5'-CC CCC CCC AAG CTT CTA GTT GAG GTT GGG AAC CAC GTC-3'; MtTPx Reverse: 5'-CCC CCC AAG CTT CTA GGC GCC CAG CGC GGC-3'). The PCR products were cloned into cloned pET22b(+) digested with corresponding set of enzymes. Expression and purification was done as described for MtAhpC.

Example 2
Measurement of NAD(P)H consumption

MtMycR assay

The activity of AhpC with MycSH and MycR was tested in a system that contained 450 μM NADH or NADPH respectively, 0.2 μM - 20 μM MtMycR, 0 μM - 20 μM MycSH, 0.2 μM - 20 μM MtAhpC, 73 μM t-boOH in 50 mM HEPES, 1 mM EDTA pH 7.4 in a final volume of 500 μl. The decrease in A₃₄₀ was monitored over time.

MtAhpD/MtAhpC assay

The specific activity of MtAhpC with MtAhpD was obtained by measuring initial velocities monitoring the NADH oxidation at 340 nm at 25°C. The test system contained 450 μM NADH, 200 nM MtAhpD, 0.4 U bovine liponamide dehydrogenase (Sigma, Taufenkirchen, Germany), 50 μM liponamide, 0.5 μM - 10 μM MtAhpC, 73 μM t-boOH in 50 mM HEPES, 1 mM EDTA pH 7.4 in a final volume of 500 μl. The reaction was preincubated for 15 min at 25°C and started by addition of the peroxide. For the AhpD-
dependent kinetic studies of MtAhpC the conditions were nearly the same as for the measurement of the activity using 5 μM MtAhpC and fixed MtAhpD concentrations. A range between 5 μM – 65 μM t-bOOH was used for the calculation of the kinetic pattern.

MtAhpC and MtTPx assay

The specific activities of MtAhpC with MtTrxC and MtTPx with MtTrxB or MtTrxC respectively were obtained by measuring initial velocities monitoring the NADPH oxidation at 340 nm at 25°C. The test system contained 450 μM NADPH, 10 μM MtTR, 10 μM MtTrxB or MtTrxC, 0.5 μM – 15 μM MtAhpC or 0.2 μM – 5 μM MtTPx, 100 nM bovine catalase (Roche, Mannheim, Germany), 73 μM t-bOOH in 50 mM HEPES, 1 mM EDTA pH 7.4 in a final volume of 500 μl. The reaction was preincubated for 15 min at 25°C and started by addition of the peroxide. MtTR and MtTrx alone showed NADPH consumption. It was reported, that thioredoxin reductases produce H₂O₂. In this test system this would start the redox cascade. To reduce this basal acitivity to a minimum, catalase was added. For the kinetic studies of MtAhpC and MtTPx the conditions were nearly the same as for the measurement of the activities using 5 μM of the peroxidase and fixed thioredoxin concentrations. A range between 5 μM – 65 μM t-bOOH was used for the calculation of the kinetic pattern. Specificity for H₂O₂, cumene hydroperoxide, linoleic acid hydroperoxide and phosphatidyl cholin hydroperoxide, both prepared from soybean lipoxygenase-catalyzed oxidation of the corresponding lipids, were investigated accordingly.
Example 3

Steady-state-kinetics

Substrate turnover by peroxidases was monitored continuously by NAD(P)H consumption in a coupled test system. The data was analyzed according to Dalziel, (1957), Acta Chem. Scand. 11, 1706-1723. Results are presented in terms of the general Dalziel-equation for two substrate reactions, in which the term $\Phi_{1,2}/[A][B]$ is zero ($[E_0]/v = \Phi_0 + \Phi_1/[A] + \Phi_2/[B] + \Phi_{1,2}/[A][B]$; see table 1). In this equation $[E_0]$ means total enzyme molarity, [A] and [B] are concentrations of the oxidizing, and reducing substrate, respectively. The empirical coefficients $\Phi_0$, $\Phi_1$, $\Phi_2$ and $\Phi_{1,2}$ are functional characteristics of the particular enzyme. The reciprocal net forward rate constant $k'_1$ by definition equals $\Phi_1$ and $\Phi_2$ describes a reductant-dependent step, thus must be the reciprocal $k'_2$. By analogy, $\Phi_0$ is the reciprocal $k_{cat}$, which can be any of the rate constants determining the a substrate-independent step.

Example 4

Gel permeation of native MtTPx

MtTPx was reduced by adding 200 mM DTT and chromatographed on a Superdex 75 column (Amersham Biosciences, Freiburg, Germany). Protein molecular weight standards (Serva, Heidelberg, Germany) were used for column calibration. Eluted peroxidase was identified by activity measurements and SDS-PAGE. For size determination of oxidized MtTPx, the enzyme was incubated with
300 µM t-bOOH at room temperature for 30 min. The molecular weight was then estimated by gel filtration chromatography as described above.

**Summary of examples**

Figure 1 shows the oxidative activation of MtAhpC and MtAhpD by t-bOOH. MtAhpC activity with MtAhpD as cosubstrate was tested in the conventional test system (11). Trace A: The reaction was preincubated for 15 min at 25°C and started by the addition of t-bOOH. Trace B: The reaction except MtAhpC, MtAhpD and t-bOOH was preincubated for 15 min at 25°C. MtAhpC, MtAhpD and the hydroperoxide were preincubated together for 2 min at 25°C and the reaction was started by the addition of this mixture.

Figure 2 shows steady-state-kinetic analysis of MtTPx with MtTrxB or MtTrxC as cosubstrates. Substrate turnover by MtTPx was monitored continuously by NADPH consumption in a coupled test system at 25°C and pH 7.4 with t-butyl hydroperoxide as substrate and MtTrxB as cosubstrate. Three sets of data were analyzed according to Dalziel (Dalziel, (1957), *supra*). Results are presented in terms of the general Dalziel-equation for two-substrate reaction ([E₀]/v = Φ₀ + Φ₁/[ROOH] + Φ₂/[reductant]; see table 1).

Figures 2A is an example of a primary Dalziel plot of the MtTPx/MtTrxB system showing enzyme-normalized inverse initial velocities ([E₀]/v) in dependence of reciprocal hydroperoxide concentration at different MtTrxB concentrations that were
kept constant by continuous regeneration. The slopes correspond to $\Phi_1$.

Figure 2B is an example of a secondary Dalziel plot: The ordinate intercepts of (A), i.e., the apparent maximum velocities at $[A] = \infty$, are plotted against $1/[MtTrxB]$ for evaluation of Dalziel coefficient $\Phi_2$ (slope), limiting $K_{MMTTrxB}$ (abscissa intercept is $1/K_{MMTTrxB}$), and real maximum velocities (ordinate intercept is $\Phi_0 = [E_0]/V_{max}$). Primary data of three independent analyses, as shown in (A), were used to obtain means and standard deviations.

Figure 2C is an example of a primary Dalziel plot of the MtTPx/MtTrxC system showing enzyme-normalized inverse initial velocities $([E_0]/v)$ in dependence of reciprocal hydroperoxide concentration at different MtTrxC concentrations that were kept constant by continuous regeneration.

Table 1 summarizes the kinetic constants of MtAhpC and MtTPx. Rate constants were derived from steady-state analysis as shown in Figure 2; the empirical Dalziel coefficients $\Phi_0$, $\Phi_1$ and $\Phi_2$ correspond to the reciprocal values of $k_{cat}$, $k'_1$ and $k'_2$, the latter two being net forward rate constants for the reaction of the enzymes with $t$-bOOH and the reductants, respectively.
Table 1

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<th>donor substrate</th>
<th>$k_{cat}$ [s$^{-1}$]</th>
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<th>$K_{M \text{ ROOH}}$ [μM]</th>
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<td>MtAhpC</td>
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<td>MtTPx</td>
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Table 2 summarizes the specificity of MtAhpC and MtTPx for hydroperoxide substrates. The activities with t-butyl hydroperoxide (t-BOOH) were set to 100%. LOOH, PCOOH and COOH mean linoleic acid hydroperoxide, phosphatidyl cholin hydroperoxide and cumene hydroperoxide, respectively. The activities with H$_2$O$_2$ were difficult to be estimated due to the basal activity of TR/Trx system.
Table 2

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Reference list

26. WO 96/19578 A2
27. WO 03/004479 A1
Claims

1. An enzyme-based assay for detecting inhibitors of the anti-oxidant defense system in *Mycobacterium* comprising
   (i) a reductant;
   (ii) a thioredoxin reductase or a peroxidase;
   (iii) a donor substrate capable of receiving a hydrogen atom from (ii); and
   (iv) a hydroperoxide,
   for determining the consumption of the reductant.

2. The assay of claim 1, wherein the thioredoxin reductase and/or the peroxidase and/or the donor substrate is an enzyme derived from *Mycobacterium*, preferably *Mycobacterium tuberculosis*.

3. The assay of claim 1 or 2, wherein the enzyme is expressed in heterologous host cells.

4. The assay of claim 3, wherein the host cells are selected from the group consisting of bacteria, yeasts, insect cells and mammalian cells, preferably cells of *Escherichia coli*, *Saccharomyces cerevisiae* and *Spodoptera frugiperda*.

5. The assay of any of the preceding claims, wherein the reductant is NADH, NADPH and/or FADH.

6. The assay of any of the preceding claims, wherein the peroxidase is a peroxiredoxin, preferably AhpC and/or TPx
of *Mycobacterium*, more preferably of *Mycobacterium tuberculosis*.

7. The assay of any of the preceding claims, wherein the donor substrate is selected from the group consisting of glutathione and thioredoxin, preferably thioredoxin B (TrxB) and/or thioredoxin C (TrxC) of *Mycobacterium*, more preferably of *Mycobacterium tuberculosis*.

8. The assay of any of the preceding claims, wherein the hydroperoxide is selected from the group consisting of hydrogen peroxide, t-butyl hydroperoxide, cumene hydroperoxide, linoleic acid hydroperoxide, phosphatidyl choline hydroperoxide and any oxidant comprising a hydroperoxy group.

9. A method for manufacturing a pharmaceutically active compound for treating and/or preventing a *Mycobacterium* associated disease, comprising the steps of
   (a) providing an enzyme-based assay of any of claims 1 to 8;
   (b) providing a test sample comprising a potentially active compound;
   (c) contacting (a) and (b);
   (d) determining consumption of a reductant;
   (e) recovering the active compound.

10. The method of claim 9, wherein the reductant is NADH, NADPH and/or FADH.
11. The method of claim 9 or 10, wherein the active compound is combined with suitable pharmaceutically acceptable fillers, excipients and/or additives.

12. The method of claim 11, wherein the pharmaceutically acceptable filler, excipients and/or additives are selected from the group consisting of physiological saline solution, demineralized water, stabilizers, proteinase inhibitors and gel formulations.

13. A use of the pharmaceutically active compound identified by the method of any of claims 9 to 12, for the preparation of a medicament for treating and/or preventing a *Mycobacterium* associated disease.

14. The use of claim 13, wherein the disease is tuberculosis.
Figure 1
Figure 2A
secondary plot

Figure 2B
primary plot

Figure 2C
SEQUENCE LISTING

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**INTERNATIONAL SEARCH REPORT**

**A. CLASSIFICATION OF SUBJECT MATTER**

IPC 7 C12Q1/18  C12Q1/26  C12Q1/28

According to International Patent Classification (IPC) or to both national classification and IPC.

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 C12Q

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, COMPENDEX, BIOSIS, PAJ, WPI Data, MEDLINE, EMBASE

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

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**Further documents are listed in the continuation of box C.**

**Patent family members are listed in annex.**

- **A** document defining the general state of the art which is not considered to be of particular relevance
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- **O** document referring to an oral disclosure, use, exhibition or other means
- **P** document published prior to the international filing date but later than the priority date claimed
- **T** later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
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- **Z** document member of the same patent family.

Date of the actual completion of the international search

28 September 2004

Date of mailing of the international search report

06/10/2004

Name and mailing address of the ISA

European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fac. (+31-70) 340-3016

Authorized officer

Tuynman, A
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