Redox modulation of thimet oligopeptidase activity by hydrogen peroxide

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Thimet oligopeptidase (EC 3.4.24.15, TOP) is a cytosolic mammalian zinc protease that can process a diversity of bioactive peptides. TOP has been pointed out as one of the main postproteasomal enzymes that process peptide antigens in the MHC class I presentation route. In the present study, we describe a fine regulation of TOP activity by hydrogen peroxide (H₂O₂). Cells from a human embryonic kidney cell line (HEK293) underwent an ischemia/reoxygenation-like condition known to increase H₂O₂ production. Immediately after reoxygenation, HEK293 cells exhibited a 32% increase in TOP activity, but no TOP activity was observed 2 h after reoxygenation. In another model, recombinant rat TOP (rTOP) was challenged by H₂O₂ produced by rat liver mitoplasts (RLMt) alone, and in combination with antimycin A, succinate, and antimycin A plus succinate. In these conditions, rTOP activity increased 17, 30, 32 and 38%, respectively. Determination of H₂O₂ concentration generated in reoxygenated cells and mitoplasts suggested a possible modulation of rTOP activity dependent on the concentration of H₂O₂. The measure of pure rTOP activity as a function of H₂O₂ concentration corroborated this hypothesis. The data fitted to an asymmetrical bell-shaped curve in which the optimal activating H₂O₂ concentration was 1.2 nM, and the maximal inhibition (75% about the control) was 1 μM. Contrary to the oxidation produced by aging associated with enzyme oligomerization and inhibition, H₂O₂ oxidation produced sulfenic acid and maintained rTOP in the monomeric form. Consistent with the involvement of rTOP in a signaling redox cascade, the H₂O₂-oxidized rTOP reacted with dimeric thioredoxin-1 (TRx-1) and remained covalently bound to one subunit of TRx-1.

Abbreviations
Abz, o-aminobenzoyl; EDDnp, N-[2,4-dinitrophenyl]ethylenediamine; JA-2, N-[1-(R,S)-carboxy-3-phenylpropyl]Ala-Aib-Tyr-p-aminobenzoate; RLMt, rat liver mitoplasts; TOP, thimet oligopeptidase; TRx-1, thioredoxin-1.
Thimet oligopeptidase (TOP, 1.3.4.24.15) is a 77-kDa zinc metalloendopeptidase that presents the HEXXH motif in the active site [1–4]. TOP is present in a diversity of mammalian tissues. The highest levels of TOP expression have been reported to occur in the brain, pituitary gland, and testis [2,5,6]. In cells, according to the cell type, TOP has been identified in the cytosol, membranes, and nucleus, which is consistent with a diversity of physiological roles [2,7]. TOP hydrolyzes many bioactive peptides of the central nervous system and other tissues [6] including bradykinin (BK), neurotensin (NT), opioid peptides, angiotensin I and gonadotrophin-releasing hormone [8–11]. TOP also cleaves several postproteasomal peptides [12], resulting from the proteolytic activity against endogenous oxidized proteins (self-antigens) as well as on foreign antigens (from bacteria and viruses). Therefore, TOP inhibition favors the cell-surface presentation of MHC-I (major histocompatibility complex-I), in a mechanism that has been described to be dependent on ROS (reactive oxygen species) production [13–15]. TOP has also been demonstrated to handle processing of the melanoma antigen PRAME peptide in a proteasome-independent mechanism [16]. Recently, TOP expression was positively correlated with tumor malignancy both in hepatocellular carcinoma and lung cancer, making its expression a good predictive prognostic factor for the tumor evolution [17,18]. TOP activity is related to increased degradation of the Aβ peptide, the component of amyloid plaques in Alzheimer’s disease produced by the degradation of the APP (amyloid precursor protein) [8,19]. The regulation of TOP activity occurs at the level of expression and by posttranslational modification such as phosphorylation [20,21]. Another mechanism postulated as contributing to TOP regulation is a multimerization by the formation of intermolecular disulfide bonds. TOP multimerization was described to occur in vitro, in the absence of reductant [3,22], and also in cellular assays due to the glutathionylation of the Cys246 and Cys175 [23,24]. In fact, a distinguishing feature of this peptidase is its activation by reducing agents, from which the denomination thimet (thiol-metallo) derives. TOP contains 15 cysteine residues that do not form intramolecular disulfide bonds. Of these 15 residues, six are surface exposed Cys that can form interchain disulfide bonds. Among the six surface exposed Cys residues, the cluster of cysteines 246, 248 and 253 (Cys246/248/253) are the most important residues for TOP oligomerization–inactivation [3]. We identified the modulation of the TOP activity in HEK293 cells by changes in H2O2 concentrations after exposure to a hypoxia/reoxygenation treatment. Considering the well-known signaling role of H2O2 we investigate the redox modulation of TOP activity by H2O2. Beyond this TOP oligomerization–inactivation process, we could demonstrate that H2O2 can reversibly oxidize TOP thiols to sulfinic and sulfonic acids, which in turn can be recycled by regulatory reductases, modulating the activity of monomeric TOP.

Results

TOP activity in HEK293 cells increases immediately after 12-h hypoxia and decreases after 2-h reoxygenation

Human embryonic kidney cells (HEK293 cell line) were submitted to an ischemia/reperfusion conditioning that leads to H2O2 production. TOP specific activity was determined using the substrate Abz-GFSPFRQ-EDDnp and the inhibitor JA-2 in extracts of treated (12-h hypoxia or 12-h hypoxia followed by 2-h normoxia/reoxygenation) and control cells (12-h or 14-h normoxia). It is important to note that to access the cells just after hypoxia an immediate reoxygenation also occurs, because the transfer from the hypoxia chamber to the regular incubator (normoxia condition), where the cells were immediately processed. Figure 1A shows that after hypoxia HEK293 cells exhibited a mean of 32% increase of TOP activity, despite the fact that on opposite way, both TOP mRNA detected by RT-qPCR (Fig. 1B) as well as the TOP protein level detected with anti-TOP antibody in western blot analysis (inset of Fig. 1A) decreased. TOP activity determined for samples of HEK293 cells after 2 h of reoxygenation revealed a mean of 90% decrease of the enzyme activity. The TOP activity decrease detected in HEK293 cells after 2-h reoxygenation did not have the contribution enzyme expression as no difference was detected between the samples 12-h hypoxia and 12-h hypoxia/2-h reoxygenation by RT-qPCR as well by western blots with anti-TOP antibody (data not shown). Also, HEK293 cells submitted to hypoxia followed by 2-h reoxygenation produced 1 μM of H2O2 measured by Amplex Red assay. This result suggests the association of TOP activity modulation in HEK293 cells with the H2O2 production. Furthermore, considering that other factors related to hypoxia and reoxygenation might respond for the TOP modulation, we first pursued in the analysis of the direct effect of H2O2 on TOP activity, using isolated rat liver mitoplasts (RLMt) and a solution of purified rat TOP (rTOP).
H₂O₂ generated by poisoned isolated rat liver mitoplasts increases rTOP activity

The *ex vivo* approach consisted of the addition of rTOP to a suspension of rat liver mitoplasts in the absence and the presence of antimycin A, succinate, or antimycin A plus succinate. Antimycin A promotes inhibition of the complex III of the respiratory chain leading to electron escape from complex III and formation of superoxide ion. The addition of succinate, which contributes to maintaining complex III reduced, exacerbated the formation of superoxide ion. Superoxide ion produced by RLMt (rat liver mitoplasts) further disproportionated to H₂O₂ by the action of manganese superoxide dismutase [25]. Figure 2 shows a basal proteolytic activity of RLMt suspension (1 mg protein/mL) that might be assigned to mitochondrial neurolysin. The activity of rTOP solution increased more than 50% in the presence 1 mg protein/mL RLMt suspension. The increase of rTOP activity was not a sum of independent RLMt and rTOP activities. It is not possible to estimate the contribution of neurolysin activity in this condition. However, it is expected that a predominant contribution of rTOP activity in the system liver RLMt/rTOP due to competition with neurolysin activity. Furthermore, even subtracting the basal contribution of the independent RLMt proteolytic activity, an increase of ~20% rTOP activity in the presence of RLMt persisted. The addition of antimycin A or succinate, both responsible for an increase in electron escape from the respiratory chain, led to a ~80% increase in the rTOP activity.

The increase of rTOP activity in the presence of antimycin A or succinate should be ~50% by subtracting the basal contribution of RLMt. In the simultaneous presence of succinate and antimycin A, an increase of ~100% in rTOP activity was observed that should be estimated around 80% increased by subtracting RLMt basal activity.

Considering the results obtained in cells (*in vivo*) and with RLMt (*ex vivo*), it was investigated the direct effect of different concentrations of H₂O₂ on rTOP activity.
H₂O₂ concentration can modulate rTOP activity

Hydrolysis of the substrate Abz-GFSPFRQ-EDDnp by 0.1 nmol-L⁻¹ of rTOP determined in the absence and the presence of different H₂O₂ concentrations are shown in Fig. 3. The maximal activation, an increase of 42% about the control activity appeared at 1.2 nmol-L⁻¹. A progressive inhibition of the activity (15 to 75%, respectively) appeared at the concentration range of 5 to 1000 nmol-L⁻¹ of H₂O₂ added to the medium (Fig. 3). The curve of the modulation of rTOP activity as a function of H₂O₂ concentration in vitro reinforced that endogenously produced H₂O₂ is responsible for the rTOP activation observed in the in vivo and ex vivo assays. The peroxide modulation of rTOP activity is specific for H₂O₂ as t-butyl hydroperoxide (t-BuOOH) did not affect significantly rTOP activity (inset Fig. 3).

H₂O₂ oxidizes TOP leading to higher oxidative states of cysteine residues

Considering that H₂O₂ modulates TOP activity in a concentration-dependent manner, a question arose: what kind of oxidative changes cause TOP activation and inhibition? In the native TOP structure, the internal cysteine residues do not form intramolecular disulfide bonds and in aged and air-oxidized TOP the external cysteine residues establish interchain disulfide bonds leading to TOP oligomerization and inhibition [3,22]. In the case of H₂O₂-oxidized rTOP, the SDS/PAGE demonstrated that even in inhibitory concentrations, H₂O₂ did not promote rTOP oligomerization (Fig. 4A,B).
Figure 4A shows that the bands of monomeric, dimeric and trimeric rTOP did not present significant changes after the treatment with different H$_2$O$_2$ concentrations. The incapacity of H$_2$O$_2$ to promote rTOP oligomerization was corroborated by the data of monomer/oligomer ratio obtained by densitometry (Fig. 4B). Therefore, the redox state of cysteine residues was investigated using the reagent NBD-Cl (7-nitrobenz-2-oxa-1,3-diazole chloride) that has a specific absorbance spectrum peaking at 347 nm when linked to sulfenic acid. Figure 4C, thin solid line, shows that the reaction of air-oxidized native rTOP with NBD-Cl yielded a sample with an absorbance spectrum peaking at 420 nm assigned to rTOP-S-NBD. This spectrum did not exhibit bands in the 300- to 350-nm region. When rTOP aged and oxidized in the air was reduced by the reagent TCEP (tris(2-carboxyethyl)phosphine), a significant increase of the band peaking at 420 nm was observed (Fig. 4C, thick solid line). That is consistent with the conversion of cysteine residues involved in the inter-chain disulfide bonds to reduced cysteine with –SH groups that are available for the formation of rTOP-S-NBD adducts. The sample of rTOP treated with 100 µM H$_2$O$_2$ exhibited a decrease in the absorbance at 420 nm concomitant with the increase of absorbance at 320–350 nm (Fig. 4C, gray line). These results are consistent with the formation of a rTOP-SOH, the first step could involve nucleophilic attack of the rTOP sulfenic acid by a thiolate of a subunit of dimeric TRx (step a). The structural change of TRx promoted by the ligation of TOP could lead to a reaction of interchain/intrachain disulfide exchange (step b) with consequent formation of a rTOP-TRx covalent complex and the liberation of a TRx monomer. Another possibility is that in the putative noncovalent complex of dimeric TRx and rTOP-SOH, a cysteine residue of rTOP can act as a resolving cysteine leading to the formation of a disulfide bond (step a‘ and b‘). The rTOP disulfide bond should be a target for the nucleophilic attack of a TRx1 thiolate followed by the occurrence of an inter-chain/intrachain disulfide exchange (step c). In cells, the formation of TRx-rTOP complex could be reduced by thioredoxin reductase (TRxR). The latter mechanism is similar to that described for the reduction of peroxiredoxin (Prx) by TRx1 [27]. TRx1 uses the free thiols of Cys32 and Cys35 to catalyze the selective reduction of disulfide bonds of target proteins. However, TRx1 can also reduce cysteine sulfenic acids probably using the same catalytic residues, that is Cys32 and Cys35 [28]. The possible conversion of dimeric TRx1 to the monomeric form raises the possibility of further redox regulation processes that deserves future investigations.
Activity of TOP C246S, C248S, and C253S triple mutant is not affected by H2O2

The cysteine residues Cys246/248/253 are important in the rTOP oligomerization–inactivation process because these residues take part in intermolecular disulfide bonds [3]. Therefore, we investigated the participation of these residues in the modulation of rTOP activity by H2O2. For this purpose, a rTOP mutant with these three cysteine residues replaced by serine residues was used. Figure 7A shows that triple-mutated rTOP C246S, C248S, and C253S possesses only 20% of the activity of WT (wild-type) rTOP. Unlike WT rTOP (Fig. 7B), the activity of the triple-mutated rTOP is not affected by DTT (1,4-dithiothreitol; Fig. 7C). Similarly, H2O2 had no effect on triple-mutated rTOP (Fig. 7D). This result is consistent with the key residues Cys246, Cys248, and Cys253 as the target for the modulatory effect of H2O2 on rTOP activity. Taken together, the results suggested that modifications at the cluster formed by the Cys residues 246, 248, and 253 affect rTOP activity, even if this modification does not lead to rTOP dimerization/oligomerization. We discarded possible interferences of the H2O2 treatment in the activity assays used herein through H2O2 treated rTOP kinetic assays with the FRET substrate Abz-GFSPFRQ-EDDnp and, inhibitory assays with the JA-2 inhibitor (data not shown). The structural rearrangements of TOP that leads to allosteric regulation remain unexplained because these residues are quite distant from the enzyme active site. Nevertheless, in fact, the structural basis to explain the inactivation of dimeric TOP TOP leads to the same question.

Discussion

Hydrogen peroxide (H2O2) is a long-lived and nonradical pro-oxidant molecule initially related to oxidative damages and now also related to cell signaling by a substantial body of evidence. The recent studies demonstrating that aquaporin-3 regulates the transmembrane traffic of H2O2 in addition to the specificity for signaling targets refuted the previous common idea of H2O2 as a freely diffusible molecule in cells and tissues. In fact, a diversity of recent studies has described H2O2 stabilized in cell compartments [29]. A diversity of mechanisms produces H2O2 in different cell compartments. Electron escape from respiratory complexes in mitochondria was the early source described for H2O2 generation in cells. Xanthine oxidase, lipoxygenase, and myeloperoxidase are also important sources of the generation H2O2 [30,31]. More recently the enzymes of the Nox (NADPH oxidase) family emerged as a source of H2O2 strongly related to signaling mechanisms. The Nox family enzymes firstly described as associated with the plasma membrane of phagocytic cells are now identified in a diversity of subcellular compartments of nonphagocytic cells such as mitochondria, nucleus, and endoplasmatic reticulum. Furthermore, a single cell type can express different Nox isoforms targeted to several subcellular compartments [32,33]. Literature data have shown that cysteine residues of different proteins are likely the principal target of the signaling action of H2O2. The redox signaling on cysteine residues requires reversible oxidation of thiol group to sulfenic acid intermediates (Cys-SOH) [34]. The irreversible oxidation of cysteine residues to sulfinic and sulfonic acid has been associated with injury and cell death [35,36]. The protein tyrosine phosphatases (PTP) are well-known targets of H2O2 generated by Nox [37]. Another family of enzymes regulated by ROS is the family of matrix metalloproteinases (MMPs) [38]. These enzymes have their expression and activation regulated by ROS. Interestingly, in the present study, we observed a fine regulation of a cellular metalloproteinase by H2O2. TOP plays an important role in the presentation of MHC-I by cells. The degradation of irreversibly
oxidized proteins by proteasomes generates a diversity of peptides that are almost all digested by cytosolic peptidases (Fig. 8). The peptides that escape from the degradation in the cytosol are translocated by the TAP (transporter associated with antigen processing) complex from the cytosol into the endoplasmic reticulum (ER). In the ER, the antigenic peptides bind the major histocompatibility complex class I (MHC class I) and are transported to the cell surface to be recognized by antibodies on the surface of lymphocyte T

![Fig. 8. Regulation of TOP activity by H$_2$O$_2$ concentration in cells.](image-url)
TOP activity control by H₂O₂

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Unlike the cell proteins responsible for the production of antigenic peptides, TOP is not regulated by interferon-γ. Therefore, considering the probable significant participation of TOP in the presentation of MHC class I at the cell surface and other signaling functions not yet elucidated it is reasonable to infer that this enzyme has a regulation mechanism. The synthesis of TOP occurs in cytosolic ribosomes without pre or pro inactive forms and further, no endogenous TOP inhibitors have been identified. Therefore, the regulation of TOP has been searched for at the transcriptional level [20,21]. On the other hand, considering the high thiol content of TOP, it is also reasonable to infer a regulation for this enzyme by a thiol-centered redox mechanism. The results presented here showing a short-term activation and long-term inhibition of TOP in cells submitted to a hypoxia/reoxygenation model (in vivo assay), an activation of TOP in the presence of RLMt stimulated to increase H₂O₂ production (ex vivo assay) and TOP activation by nanomolar concentrations of H₂O₂ support H₂O₂ as the key molecule for the control of TOP activity control. Therefore, the present study brings to light the following questions about TOP: (a) What is the role of the high content of cysteine residues as they do not participate in the catalytic process? (b) How can a cytosolic enzyme acting in a reducing microenvironment be regulated by dimerization stabilized by disulfide bonds? (c) How does dimerization via disulfide bonds correlate with a probable control of TOP at the gene expression level? At least, Cys246, Cys248, and Cys253 seem to be involved in redox regulation of TOP. The present results by linking TOP regulation with a H₂O₂ concentration in cells suggest that the formation of sulfenic, sulfenic and sulfonic acids in addition to the interchain disulfide bonds are responsible for the control of the metalloproteinase activity. To identify the presence of higher forms of oxidation on TOP, we used the NBD trap, which maximally absorbs near 337 nm when attached to SOH as S-O-NBD. When H₂O₂ mildly oxidizes TOP, we note an increase in this band. When we overoxidized rTOP, this absorption decreased, suggesting that higher oxidized forms were generated as NBD is unable to react with these modifications. When tested against Cys mutants both oxidative and reducing profiles were abrogated. The peculiar structure of TOP, bearing 15 cysteine residues some of them surrounded by acidic and basic residues, makes this enzyme a candidate to be a H₂O₂ sensor. TOP has the majority of its Cys residues located on the internal side of the structure but in an arrangement unfavorable to form internal disulfide bonds. However, higher oxidative states of cysteine may occur regarding the generation of sulfur–oxygen bonds. These states are the unstable (naturally reversible) sulfenic acid (SOH) that is formed by low levels of H₂O₂ in cells and the sulfenic (SO₂) and sulfonic (SO₃) acids, the two latter are associated with conditions of oxidative stress with high production of H₂O₂ and naturally irreversible. These cysteine derivatives are produced through the reduction of H₂O₂ by the low pKₐ cysteine residues of key proteins. Subsequently, they are recycled to the SH form by the action of specific enzymes in events that composes the signaling pathway of H₂O₂. We discard the effect of TOP oligomerization in H₂O₂-treated TOP by following SDS/PAGE of monomeric TOP. TOP oligomerization was reported in HEK293 challenged by 100–400 μM of H₂O₂ [23]. H₂O₂ is diffusible across membranes, but studies of Choi et al. [39] demonstrated that exogenously added H₂O₂ has been shown to be less effective for triggering signaling responses than endogenously produced H₂O₂. In fact, the mechanisms of H₂O₂ deactivation present in cells suggest that the signals charged in this molecule are only transmitted over relatively short distances [40]. The present results obtained in vivo, ex vivo and in vitro strongly raises the possibility that the action of this peptidase can be modulated by the redox state of cells. Higher oxidative forms of cysteine are becoming important physiologically and postulated as a chemical modification that links protein function to the oxidative status of cells. For example, the formation of a stabilized cysteine sulfenic acid in the DJ-1 protein is critical for the mitochondrial function of this protein associated with inherited Parkinsonism [41]. Thus, our results suggested that modifications at the cluster formed by the Cys residues C246, C248, or C253 affects TOP activity, even if this modification does not lead to TOP dimerization/oligomerization. The exact structural rearrangements that occur in the enzyme that explains this allosteric regulation remains to be elucidated. The decrease of TOP expression in response to the short-term activation of the TOP cell content after reoxygenation links the chemical and structural modulation of TOP activity with the regulation of the activity of this enzyme at the level of gene expression. Figure 8 illustrates a possible mechanism to regulate MHC class I presentation via inhibition or activation of TOP according to the concentration of H₂O₂ in cells. H₂O₂ generated by an exogenous source such as neutrophils or by endogenous sources such as electron escape from respiratory chain and enzymes of the Nox family promotes direct or indirect oxidation of proteins. Proteasomes degrade the oxidized proteins generating peptides. These peptides could have a
diversity of signaling functions including the participation in the MHC class I. In conditions in which the
increase of H$_2$O$_2$ in the cell is relatively low, TOP is
activated, promotes the proteolysis of the peptides and
impairs the presentation in the MHC class I. Other-
wise, when the H$_2$O$_2$ concentration is relatively higher,
TOP is inhibited, and the peptides follow the secre-
tory pathway. In the former condition, that is when the
H$_2$O$_2$ concentration increases slightly in cells, TOP
oxidation results in the formation of sulfenic acid and the enzyme activation could be reverted by
TRx/TRxR (thioredoxin reductase) system. On the
other hand, TOP irreversibly oxidized by high con-
centrations of H$_2$O$_2$ could be directed to proteolysis
in the proteasome. It is important to note that H$_2$O$_2$
produced in cells submitted to hypoxia and reoxy-
genation could be produced by the electron escape
from the respiratory chain as well as from Nox activ-
ity in different cell compartments. Hereafter, other
studies must be conducted to elucidate the most sig-
ificant source of H$_2$O$_2$ that can modulate TOP
activity.

The relationship of the cysteine-rich structure of
TOP with its biological role and regulation has been a
challenge, particularly the unfeasible regulation by
enzyme oligomerization. Taken together, the results
presented here demonstrate for the first time that
TOP, a thiol-rich enzyme, can potentially participate
in cell signaling mediated by H$_2$O$_2$ with participation
of thioredoxin. The conclusion is supported by the fine
oxidative regulation of TOP promoted by H$_2$O$_2$-
duced formation of sulfenic acid that was demon-
strated in vivo, ex vivo, and in vitro assays.

Materials and methods

Cell culture

HEK293 cell line was cultured in DMEM (Vitrocell, São
Paulo, Brazil) containing 10 µg·mL$^{-1}$ of streptomycin
(Vitrocell) and supplemented with 10% fetal bovine serum
(Vitrocell). HEK293_TOP cells which overexpress TOP
protein were previously generated [42] and cultured under
the same conditions.

Hypoxia/reoxygenation assay

Cells were seeded in triplicate, at a concentration of 7 \times 10^6
cells (70% of confluence) in T-75 bottles (Corning Incor-
porated, Corning, NY, USA), in 10 mL culture medium. The
cultured cells were grown for 24 h before the experiments at
37 °C in a humidified 5% CO$_2$ incubator. Subsequently, the
cell culture medium was removed, and the cells were washed
once with PBS. In the following, cells were incubated in two
different conditions: 12 h in the normal atmosphere (control)
and 12 h in a hypoxia chamber under 5% O$_2$ atmosphere
[43]. A model of ischemia and reperfusion consisted in main-
taining cultured for 12 h in a hypoxic chamber with subse-
quent return to the normal atmosphere. In this chamber, the
atmosphere composition was 5% O$_2$, 5% CO$_2$, and 90% N$_2$,
followed by a 2-h incubation in a regular incubator with
atmosphere composition of 30% O$_2$, 5% CO$_2$, and 65% N$_2$.
TOP activity was measured immediately after the removal of
the hypoxia chamber condition and after 2 h of incubation
under a normal atmosphere.

Rat liver mitoplasts

Mitochondria were isolated by conventional differential
centrifugation from the livers of adult Wistar rats weight-
ing approximately 180 g and receiving food and water
ad libitum in a light-controlled room (12 h light/dark
cycles). Each liver was reached through a bilateral
abdominal incision. All procedures here were conducted
according to an approved institutional animal experimen-
tation protocol (committee of ethical research approval
number 0540/11), which follows the ‘Guide for the Care
and Use of Laboratory Animals in Research’ (in accord-
dance with the National Institutes of Health, USA). The
homogenate was prepared in 250 mM sucrose, 1.0 mM
EGTA, and 5.0 mM HEPES buffer (pH 7.2). The mito-
ochondrial suspension was washed twice in the same
medium containing 0.1 mM EGTA, and the final pellet was
diluted in 250 mM sucrose to a protein concentration of
80–100 mg·mL$^{-1}$ [44,45]. RLMt (mitochondria devoid of
the outer membrane) were prepared as described by Ped-
eresen et al. [46–48]. Rat liver mitoplasts were incubated
in 50 mM Tris buffer, 100 mM NaCl, pH 7.4, at 37 °C.
When indicated, the medium contained 2.5 mM succinate
and 5 µm antimycin A. For every specified condition,
rTOP was added to the RLMt-containing medium and the
TOP specific enzymatic activity determined using the
fluorescent substrate Abz-GFSPFRQ-Eddnp (10 µm) [49]
and the inhibitor JA-2 (10 µm) [50,51].

Quantification of H$_2$O$_2$ produced by RLMt

Production of H$_2$O$_2$ was determined in 0.125 or 0.25 mg
protein·mL$^{-1}$ of mitochondrial suspensions in 10 mM KCl,
2 mM HEPES pH 7.4 [52]. The production of H$_2$O$_2$ was
stimulated by the presence of 2.5 mM succinate, and 5 µm
antimycin A. H$_2$O$_2$ was quantified by Amplex Red (Sigma-
Aldrich-Merck, Darmstadt, Germany) in the concentration
of 25 µM [53,54]. Briefly, Amplex Red is oxidized to resoru-
fin by 0.5 U·mL$^{-1}$ horseradish peroxidase (HRP) added to
the medium. The activation of HRP is dependent on H$_2$O$_2$
produced by RLMt. Resorufin is a fluorescent derivative
detected by a spectrofluorimeter operating at 563 nm for
excitation and 587 nm for emission. This technique displays good signal/noise ratios and little artificial interference [55,56]. Indeed, controls conducted in the absence of mitochondria or the absence of peroxidase indicate that nonspecific probe oxidation is negligible (<1% of the increment observed in the presence of mitochondria and peroxidase). Also, fluorescence increments are largely suppressed (>90%) in the presence of added catalase, indicating the response is mostly due to H₂O₂ production. Calibration was conducted by adding H₂O₂ at known concentrations (ε₂₄₀ nm = 43.6 M⁻¹·cm⁻¹).

Expression of recombinant wild-type and mutant rat TOP

Clones construction, mutant generation, expression, and purification were performed as already published with some modifications [57]. Briefly, recombinants wild-type and mutant rTOP were expressed in Escherichia coli BL21 (DE3) strain (Merck-Novagen, Darmstadt, Germany) as a poly(His) tag fusion protein using the expression vector pET28a(+) (Merck-Novagen). Protein purification was performed by affinity chromatography. The purity of the protein was determined by Coomassie brilliant blue staining on 12% SDS/PAGE. rTOP homogeneity was larger than 95% (data not shown). Protein samples were stored at −80 °C in small aliquots.

Enzyme reduction

Preparations of purified recombinant rTOP were treated with the sulfhydryl reductant TCEP (1–10 mM) [58]. After incubation, rTOP preparations were passed through a PD-10 desalting column (GE Healthcare Life Sciences, Buckinghamshire, UK) to remove the reducing agents completely. After determination of protein concentration by absorbance (ε₂₈₀ nm=78,115 M⁻¹·cm⁻¹), aliquots of reduced rTOP preparations were taken for further incubation, hydrolytic assays, determination of TOP-reduced Cys residues.

Determination of rTOP-reduced Cys residues

These assays were carried out as previously described [59]. Briefly, 150–250 µg of rTOP was resuspended in 300 µL of 30 mM Tris (pH 7.4) containing 8 M urea. After complete dissolution, samples were taken for reading at 280 nm. Afterward, 10 µM DTNB (final concentration, Sigma-Aldrich-Merck) was added to the samples and incubated for 40 min in the dark followed by reading at 412 nm. Protein concentration was calculated as described below. The concentration of the CyS–TNB complex was calculated from the ε₄₁₂ nm=14,150 M⁻¹·cm⁻¹. The number of reduced Cys residues was calculated by the molar ratio protein/CyS–TNB complex.

SDS/PAGE analysis

Electrophoresis was performed on 12% polyacrylamide gels containing sodium dodecyl sulfate [60]. The samples were prepared under reducing and non-reducing conditions, depending on the presence of β-mercaptoethanol in the sample buffer. Proteins were visualized with Coomassie Blue. All reactions were assayed at room temperature (25 °C) using Tris buffer pH 7.4 (Tris 50 mM, NaCl 100 mM). The reactions were stopped by ice-cold freezing. 1–2 µM of thimet oligopeptidase were incubated with increasing concentrations of a reactant or fixed concentration. The SDS/PAGE gels were scanned, and the densitometry of protein bands on gel images was performed using the software IMAGEJ [61]. The acquired data were further analyzed using GRAFIT software (v.5.0, ERITHACUS Software) [62].

Western blot

For the western blot, 1 µg of recombinant TOP and 30 µg of cell extract proteins per lane were separated by 12% SDS/PAGE and electrotransferred to a PVDF membrane Hybond-P (GE Healthcare) and then blotted with primary antibodies according to the manufacturer’s instructions anti-24.15 (anti-THOP, Proteimax Biotechnology, São Paulo, Brazil) or anti-GAPDH (Proteimax Biotechnology) detected by an anti-rabbit secondary antibody (T2767, Invitrogen Corporation, Carlsbad, CA, USA). Data were analyzed by western blot densitometry using IMAGEJ software [61].

Quantitative PCR

For the quantitative real time, RNA was isolated from cell plates using TRIzol Reagent (Thermo Fisher Scientific Inc., Waltham, MA, USA), according to the manufacturer’s instructions. Two micrograms of total RNA were treated with DNase (Ambion Inc., Austin, TX, USA) and reverse-transcribed into cDNA using Superscript III transcriptase and random primers (Invitrogen Corp.), according to the manufacturer’s instructions. An aliquot of the cDNA (10 ng) was used as template for qPCR amplification. qPCRs were carried out in a 12-µL PCR containing 3.2 pmol of each specific primer, 1x SYBR Green Master Mix (Thermo Fisher Scientific Inc.), and DEPC water. The PCR running parameter was the standard program in the ABI Prism 7500 sequence detection system (Thermo Fisher Scientific Inc.). The ΔΔCt was calculated relative to control samples and GAPDH reference gene using the comparative cycle threshold (Ct) method (2^ΔΔCt), where ΔΔCt = ΔCt test sample – ΔCt control samples and ΔCt = Ct test sample – Ct reference gene) following the MIQE guidelines [63]. The following qPCR primers were used: TOP Fw 5’-CTCCCCAGAGAGACTCAG-3’, TOP Rev 5’-GTCGTTGCTCCTGTTGGTT-3’, GAPDH Fw 5’-
ACACAGTCCATGCCCATAC-3' and GAPDH Rv 5'-TCCACCCACGTGCTGTA-3' (IDT, Integrated DNA Technologies).

Enzyme activity assay

The hydrolysis of the fluorogenic substrates Abz-GFSPFRQ-Eddnp was performed as previously described [64,65] in Tris 50 mM, NaCl 100 mM buffer, pH 7.4, at 37 °C. In the following, we measured the fluorescence at λem = 420 nm/λex = 320 nm in a spectrofluorometer (F-7000, Hitachi, Tokyo, Japan). The plate with 1 cm path length, containing 300 μL of the substrate solution was placed in a thermostatically controlled cell compartment for 5 min. In the following, the enzyme solution was added to the medium and the kinetic of fluorescence recorded for 5–10 min. The data were fitted, and the slope converted into Moles of hydrolyzed substrate per minute based on the fluorescence curves of standard peptidase solutions before and after total enzymatic hydrolysis. The concentration of the peptide solutions was calculated from the colorimetric determination of the 2,4-dinitrophenyl group (ε365 nm = 17,300 M⁻¹·cm⁻¹). The enzyme concentration for initial rate determination was chosen for hydrolyzing less than 5% of the substrate present. All the obtained data were fitted by nonlinear least squares equations, using GRAFIT software (v.5.0, ERITHACUS Software) [62].

Trapping of sulfenic acid with NBD-CI

Protein (10 μM) was exposed to oxidants (90 μM) for 30 min. NBD-CI (Sigma-Aldrich, 100 μM) was added to the samples and was incubated for 30 min, at 37 °C, after the samples were subjected to spin column (Amicon Ultra 10 kDa, Millipore, Ireland). Electronic absorption spectra were carried out in a photodiode Multispe 1501 spectrophotometer (Shimadzu Scientific Instruments, Columbia, MD), using quartz cuvettes with 1.0- and 0.1-cm optical path and 0.5-nm slit [66].

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Author contributions

MYI and JCF designed and performed the assays, analyzed the data, and wrote the manuscript. CHY, LVB, AM, and JMG performed the assays and analyzed the data. VO and ILN designed the work, analyzed the data, and wrote the manuscript.

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