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# Reversible Oxidation of the Membrane Distal Domain of Receptor PTP $\alpha$ Is Mediated by a Cyclic Sulfenamide<sup>†</sup>

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Received July 31, 2006; Revised Manuscript Received October 25, 2006

ABSTRACT: Protein tyrosine phosphatases (PTPs) are fundamental to the regulation of cellular signalling cascades triggered by protein tyrosine kinases. Most receptor-like PTPs (RPTPs) comprise two tandem PTP domains, with only the membrane proximal domains (D1) having significant phosphatase activity; the membrane distal domains (D2) display little to no catalytic activity. Intriguingly, however, many RPTP D2s share the catalytically essential Cys and Arg residues of D1s. D2 of RPTP $\alpha$  may function as a redox sensor that mediates regulation of D1 via reactive oxygen species. Oxidation of Cys723 of RPTP $\alpha$  D2 (equivalent to PTP catalytic Cys residues) stabilizes RPTP $\alpha$  dimers, induces rotational coupling, and is required for inactivation of D1 phosphatase activity. Here, we investigated the structural consequences of RPTP $\alpha$  D2 oxidation. Exposure of RPTP $\alpha$  D2 to oxidants promotes formation of a cyclic sulfenamide species, a reversibly oxidized state of Cys723, accompanied by conformational changes of the D2 catalytic site. The cyclic sulfenamide is highly resistant to terminal oxidation to sulfinic and sulfonic acids. Conformational changes associated with RPTP $\alpha$  D2 oxidation have implications for RPTP $\alpha$  quaternary structure and allosteric regulation of D1 phosphatase activity.

The reversible tyrosine phosphorylation of proteins, reciprocally controlled by tyrosine kinases and phosphatases, is crucial to the regulation of diverse cellular processes, including growth, proliferation, and differentiation (I). Redox reactions, mediated by reactive oxygen species (ROS),<sup>1</sup> control PTP activity in response to a variety of hormones and growth factors (2-6). Generation of ROS accompanies PTK activation (7, 8), with oxidation-induced inhibition of PTP activity amplifying PTK-dependent signaling (9-11).

PTP-catalyzed protein dephosphorylation proceeds via a nucleophilic displacement reaction utilizing an essential cysteine residue conserved within the PTP signature motif  $Cys(X)_5Arg$  (12-15). The low  $pK_a$  of the catalytic cysteine necessary for its function (16) renders PTPs susceptible to inactivation by ROS-induced cysteine oxidation. However,

the reversibility of redox-mediated inactivation of PTPs requires that a reversibly oxidized state of the cysteine residue be stabilized at the PTP catalytic site and that formation of SO<sub>2</sub> and SO<sub>3</sub>, two irreversibly oxidized species of cysteine, be suppressed (17). Crystallographic studies of the tyrosine specific cytosolic PTP1B revealed that the reversibly oxidized form of the enzyme was not the expected sulfenic acid or disulfide bond, but an oxidation state of cysteine termed a cyclic sulfenamide, a species not previously observed in proteins (18, 19). Generation of the cyclic sulfenamide inhibits catalytic activity by blocking the nucleophilic cysteine residue through a covalent bond of its thiol group with the neighboring main chain amide, a linkage that is accompanied by profound conformational transitions of the catalytic site PTP and pTyr loops. The oxidation properties of the catalytic Cys observed in PTP1B may be unique to the tyrosine specific subfamily of PTPs. Crystallographic and biochemical studies show that, in contrast to PTP1B, in the dual-specificity phosphatases KAP (20) and Cdc25 (21-23), lmPTPs (24), and lipid phosphatase PTEN (25), a disulfide bond linking the catalytic Cys residue to a proximal Cys is responsible for reversible oxidation and phosphatase inactivation. As in the cyclic sulfenamide of PTP1B, a disulfide bond suppresses overoxidation of the catalytic cysteine and is reversed by reductants. In another redox-regulatory mechanism, a stable and reversible sulfenic acid species was implicated as the oxidation-dependent inactive state of the dual-specificity phosphatase VHR (17).

<sup>&</sup>lt;sup>†</sup> The work in D.B.'s laboratory was funded by Cancer Research UK. The work was supported in part by the Netherlands Proteomics Centre (NPC) and grants from the research council for earth and life sciences (ALW) and ASPASIA with financial aid from the Netherlands Organisation of Scientific Research (NWO).

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<sup>&</sup>lt;sup>1</sup> Abbreviations: pNPP, p-nitrophenol phosphate; PTP, protein tyrosine phosphatase; RPTPα, receptor-like protein tyrosine phosphatase; RPTPα D1, receptor protein tyrosine phosphatase  $\alpha$  (membrane proximal) domain 1; RPTPα D2, receptor protein tyrosine phosphatase  $\alpha$  (membrane distal) domain 2; ROS, reactive oxygen species; SO<sub>2</sub>, sulfinic acid; SO<sub>3</sub>, solfonic acid; SN, cyclic sulfenamide.

The only precedent for a protein cyclic sulfenamide is the oxidized state of PTP1B, visualized directly by X-ray crystallography, and indirectly in solution by pulse-chase labeling and mass spectrometry experiments (18, 19). Interestingly, examples of cyclic sulfenamides were reported some 30 years ago in studies of the synthesis of the antibiotic cephalosporin from penicillin precursors via the nucleophilic attack of a nitrogen atom on sulfenic acids (26-28). More recently, the synthesis of a cyclic sulfenamide species in vitro from a sulfenic acid precursor was demonstrated (29). There is now abundant evidence that numerous tyrosine specific PTPs are regulated via reversible and transient oxidation, suggesting that redox regulation of these PTPs may also feature a cyclic sulfenamide species (5, 6). For example, ingel phosphatase assays demonstrate that SHP-2 was reversibly oxidized in response to PDGF signaling (30), whereas ROS have been implicated in the suppression of both PTP1B and TCPTP activities in cells stimulated with insulin (11, 31, 32) and EGF (7).

While all cytosolic PTPs comprise a single catalytic domain linked to diverse regulatory or targeting domains, the majority of transmembrane receptor-like phosphatases comprise two PTP domains arranged in tandem (1, 33). The membrane proximal D1 domain is responsible for all or most of the protein phosphatase activity with little to no phosphatase activity contributed by the D2 (distal) domain. Although essentially devoid of phosphatase activity, intriguingly most RPTP D2s share the conserved catalytic Cys and Arg residues of the PTP signature motif required for phosphatase activity. Furthermore, the crystal structures of LAR and RPTPa D2s indicated an environment of the PTP motif Cys residue essentially identical to that of catalytically active PTPs, which would be expected to promote a catalytic Cys with a reduced  $pK_a$ , enhancing its nucleophilicity (34, 35). In striking experiments, the introduction of Tyr and Asp residues at the catalytic sites of LAR and RPTPα D2, residues required for pTyr recognition and general acid catalysis in PTP1B (36), conferred tyrosine specific phosphatase activity with catalytic rates equivalent to those of RPTP D1s and cytosolic PTPs (34, 37, 38). The catalytic Cys residues of RPTP D2s are therefore as reactive and oxidation sensitive as their counterparts in RPTP D1s (39).

A role for RPTP $\alpha$  D2 as a redox sensor was suggested from the findings that  $H_2O_2$  induced numerous functional changes that were all dependent on the D2 catalytic Cys residue. Such changes include stabilization of RPTP $\alpha$  dimers, rotational coupling of the extracellular domain, and complete inactivation of D1 phosphatase activity (39, 40). Development of a generic antibody that detects oxidation of the catalytic site Cys provided a direct observation of an oxidized PTP-motif Cys residue and revealed that D2 of RPTP $\alpha$  is more sensitive to oxidation than D1 (41, 42). These results implicate a redox-dependent function of RPTP D2s, whereby the conserved Cys residue may function as a redox sensor, potentially mediating oxidation-dependent conformational changes of RPTPs and inside-out signaling (40).

Here, we investigate the structural consequences of RPTP $\alpha$ D2 oxidation by means of X-ray crystallography and explored the stability of the reversibly oxidized state in solution via catalytic activity measurements, mass spectrometry, and the oxidation specific antibody. We demonstrate that when RPTP $\alpha$ D2 crystals are exposed to oxidants ( $H_2O_2$ 

and oxidized glutathione), oxidation of the conserved Cys residue generates a highly stable cyclic sulfenamide species, identical in chemical composition to PTP1B. These studies indicate that formation of the reversible cyclic sulfenamide is a general feature of the tyrosine specific PTPs and, importantly, provide a direct demonstration of the susceptibility of the D2 catalytic Cys to oxidation, with potential implications for redox-dependent regulation of RPTP $\alpha$  conformational changes and catalytic activity.

#### EXPERIMENTAL PROCEDURES

Structure Determination. Residues Ile512-Ser787 of murine RPTPα D2 were expressed from a pET-28a expression vector (a generous gift of Joseph Noel) and purified and crystallized under conditions similar to those described previously (35) except that prior to crystallization the protein was dialyzed into 10 mM Tris-HCl (pH 8.0), 100 mM NaCl, and 2 mM DTT. Crystals were prepared using the hanging drop vapor diffusion method at 14 °C by mixing protein at 12 mg/mL with an equal volume of a reservoir solution [13– 15% (w/v) PEG 8K, 0.1 M sodium succinate (pH 5.5), 0.4 M LiSO<sub>4</sub>, and 2 mM DTT]. Crystals belong to space group C222<sub>1</sub> with two molecules per asymmetric unit and the following cell parameters: a = 74.7 Å, b = 94.7 Å, and c= 205.0 Å. This crystal form differs from that reported by Sonnenburg et al. (35) (space group P3<sub>1</sub>), although similar crystallization conditions, except for temperature, were used. Oxidation experiments were performed by incubating crystals in either 50 or 100  $\mu$ M H<sub>2</sub>O<sub>2</sub> or 10 mM oxidized glutathione for the specified times at 14 °C prior to crystal freezing (Table 1). For the reduction experiment, previously oxidized crystals were incubated in 10 mM DTT for 16 h. Data were collected at ESRF and processed using the CCP4 program suite (43). The initial structure in the series was determined by molecular replacement using the RPTPα D2 coordinates (35) (PDB entry 1P15) as a search model and refined using REFMAC, with subsequent structures being refined relative to this reference structure (Table 1).

Analysis of Oxidized PTPs Using the oxPTP Antibody. RPTPα D2 oxidation was assayed using an antibody, oxPTP, that specifically recognizes the sulfonic acid form of catalytic site cysteines of PTPs (41, 44). Briefly, the GST-RPTPa D2 fusion protein, encoding residues 504-793, was bound to glutathione—agarose beads. All incubations and washes were performed in 20 mM Tris (pH 7.5). The proteins were reduced for 30 min in 10 mM dithiothreitol (DTT). Beads were washed twice and were incubated with 0.1 or 1.0 mM H<sub>2</sub>O<sub>2</sub> for 0-24 h. Immediately after being treated, the proteins were processed for detection of (ir)reversibly oxidized catalytic cysteines by alkylation with 40 mM iodoacetic acid (IAA) for 30 min, reduction with 10 mM DTT for 30 min, and peroxidation with 0.1 mM pervanadate for 1 h (1 mM pervanadate and 1 mM Na<sub>3</sub>VO<sub>4</sub> mixed with 5 mM H<sub>2</sub>O<sub>2</sub>, final concentrations, and left for at least 5 min at room temperature) to convert reduced cysteines irreversibly into the sulfonic acid form. Between treatments, the fusion proteins were extensively washed. Different orders of treatments were used to detect the reversibly oxidized catalytic cysteine [alkylation, reduction, and pervanadate (ARP)] or the sulfinic acid form [reduction, alkylation, and pervanadate (RAP)] as outlined in Figure 4B. After processing had been carried out, Laemmli sample buffer was added, and the

Table 1: Crystallographic Data Collection and Refinement Statistics<sup>a</sup>

	50 µM H <sub>2</sub> O <sub>2</sub> for 30 min	50 μM H <sub>2</sub> O <sub>2</sub> for 1.5 h	50 μM H <sub>2</sub> O <sub>2</sub> for 3 h	$\begin{array}{c} 100\mu\mathrm{M}\;\mathrm{H_2O_2} \\ \text{for 6 h} \end{array}$	50 μM H <sub>2</sub> O <sub>2</sub> for 16 h	10 mM GSSG for 12 days	reduced DTT	DTT for 0 h, control	DTT for 16 h, control
space group	C222 <sub>1</sub>	C222 <sub>1</sub>	C222 <sub>1</sub>	C222 <sub>1</sub>	C222 <sub>1</sub>	C222 <sub>1</sub>	C222 <sub>1</sub>	C222 <sub>1</sub>	C222 <sub>1</sub>
cell parameters (Å)	a = 74.82, b = 94.79, c = 205.80	a = 75.29, b = 95.94, c = 205.84	a = 74.90, b = 94.66, c = 205.71	a = 75.0, b = 94.68, c = 205.62	a = 74.65, b = 94.79, c = 204.82	a = 75.19, b = 94.62, c = 207.39	a = 71.79, b = 97.31, c = 198.39	a = 74.63, b = 95.36, c = 204.86	a = 74.47, b = 95.39, c = 204.66
beamline	ID14.2	ID23.1	ID14.4	ID14.4	ID14.4	ID14.4	ID14.4	ID14.3	ID14.2
resolution (Å)	2.3	2.1	1.90	2.0	1.85	2.5	2.4	2.4	2.3
$R_{\text{merge}}$	0.065 (0.262)	0.082 (0.324)	0.063 (0.328)	0.055 (0.141)	0.048 (0.387)	0.076 (0.12)	0.082 (0.136)	0.109 (0.323)	0.066 (0.306)
$I/\sigma I$	7.0 (2.5)	4.8 (1.2)	6.5 (1.5)	8.4 (4.5)	9.1 (1.9)	6.1 (5.5)	5.0 (4.2)	2.4(1.0)	5.3 (1.7)
completeness (%)	99.2 (99.2)	99.4 (99.2)	95.3 (77.6)	80.3 (49.5)	74.3 (40.6)	94.2 (81.1)	94.1 (81.0)	99.9 (99.9)	99.2 (99.2)
multiplicity	3.7 (3.5)	4.0 (4.1)	3.7 (2.3)	3.1 (2.1)	4.0 (2.1)	5.0 (3.5)	4.0 (2.8)	4.0 (4.0)	4.0 (3.7)
no. of reflections	119994	176131	202962	122748	219282	128785	102476	115543	130217
detector	ADSC	ADSC	ADSC	ADSC	ADSC	ADSC	ADSC	MAR	ADSC
R-factor	0.226	0.230	0.217	0.208	0.212	0.212	0.232	0.241	0.247
$R_{\rm free}$	0.290	0.278	0.249	0.241	0.245	0.243	0.301	0.325	0.296
Cys723 MolA	60% SN <sup>b</sup>	$SN^b$	$SN^b$	$SN^b$	$SN^b$	$SN^b$	$SH^c$	$SH^c$	$SH^c$
Cys723 MolB	90% SH <sup>c</sup>	$SN^b$	$SN^b$	$SN^b$	$SO_2^d$	$SO_3^e$	$SH^c$	$SH^c$	$SH^c$

<sup>&</sup>lt;sup>a</sup> Figures in parentheses refer to the highest-resolution shell. <sup>b</sup> Cyclic sulfenamide. <sup>c</sup> Free thiol. <sup>d</sup> Sulfinic acid. <sup>e</sup> Sulfonic acid.

samples were boiled for 5 min and run on a 10% SDSpolyacrylamide gel (5 µg of protein per lane). Samples were transferred to PVDF membrane by semidry blotting. For detection with the oxPTP antibody, the blots were blocked in 0.1% BSA, 0.1% Tween in 50 mM Tris (pH 7.5), and 150 mM NaCl for 1 h or overnight and incubated with the oxPTP antibody (1.5 h at room temperature or overnight at 4 °C). Subsequently, a HRP-coupled goat anti-rabbit antibody (1:10000, BD Pharmigen) was used, and the antibody signals were visualized by enhanced chemiluminescence (ECL). The membranes were stripped and reprobed with the anti-RPTP $\alpha$ antibody (45) to monitor equal loading of the fusion protein.

MALDI-ToF Mass Spectrometry. All reactions and washing steps were performed in 20 mM Tris-HCl (pH 7.5). GST fusion proteins of RPTPα D2 were immobilized on gluthatione-Sepharose beads and incubated with 10 mM DTT. Beads were washed and incubated with 0.1 mM H<sub>2</sub>O<sub>2</sub> for 0.5-24 h at room temperature. Subsequently, beads were treated with 100 mM iodoacetamide for 30 min at room temperature, to derivatize the unaffected cysteines. Finally, beads were washed and incubated with 10 mM DTT for 10 min to reduce singly oxidized cysteines. The fusion protein was subjected to overnight tryptic digestion. For desalting, peptides were adsorbed on C18 ZipTips (Millipore) and washed with 0.1% trifluoroacetic acid (TFA). Peptides retained on the columns were eluted using α-cyano-4hydroxycinnamic acid (10 mg/mL) in 10% acetonitrile and 0.1% TFA. Peptide mass fingerprint spectra were recorded using matrix-assisted laser desorption ionization mass spectrometry (MALDI-ToF/ToF; AB 4700 proteomics analyzer, Applied Biosystems, Framingham, MA). This instrument is equipped with a 200 Hz Nd:YAG laser operating at 355 nm. Experiments were performed in a reflectron positive ion mode using delayed extraction. Typically, a total of 7500 shots were acquired per spectrum, and the signal was averaged. MALDI-ToF/ToF spectra were analyzed using the Applied Biosystems Data Explorer software.

Catalytic Activity Measurements. The catalytic activity of RPTPα D2 was measured using 50 mM p-nitrophenol phosphate (p-NPP) as a substrate in a reaction buffer of 50 mM MES (pH 6.5), 100 mM NaCl, 0.5 mM EDTA, and RPTP $\alpha$  D2 (Ile512-Ser787) at a concentration of 300  $\mu$ g/ mL. To determine the reversibility of the oxidation reaction, the activity of the RPTPa was measured (i) prior to exposure

to H<sub>2</sub>O<sub>2</sub>, (ii) following an 18 h incubation with 50  $\mu$ M H<sub>2</sub>O<sub>2</sub> at 14 °C, and (iii) after a 3 h incubation of the oxidized protein with 25 mM DTT at 20 °C. Hydrolysis of p-NPP was monitored at 410 nm, and protein concentrations were determined, after removal of the aggregated protein by highspeed centrifugation, at the three time points to estimate specific activity.

Quantitative Gel Filtration. Reduced and oxidized RPTPa D2 (Ile512-Ser787) were loaded onto an analytical gel filtration column (Superdex 200 HR10/30, GE Healthcare) at 3 mg/mL in a buffer comprising 50 mM Tris-HCl (pH 8.0) and 100 mM NaCl. The oxidized protein was prepared by dialysis against  $100 \,\mu\text{M} \,\text{H}_2\text{O}_2$  for 16 h. Oxidation of the protein was confirmed by measuring its catalytic activity. The apparent molecular weight of the protein was estimated by calibrating the gel filtration column using molecular weight standards.

### **RESULTS**

Structure of RPTPa D2. Crystallization of RPTPa D2 at 14 °C in our study yielded an orthorhombic crystal form (C222<sub>1</sub>) differing from the trigonal crystals (P3<sub>1</sub>) obtained at 4 °C by Sonnenburg et al. (35). The two molecules of RPTP $\alpha$  D2 in the  $C222_1$  asymmetric unit associate to form a dimer identical to the RPTPα D2 dimers of the P3<sub>1</sub> crystals (35). The dimer interface is created from a surface of the molecule opposing the catalytic site. Both crystallographic independent subunits of RPTPa D2 (designated MolA and MolB) adopt essentially identical conformations (rmsd of 0.6 Å for equivalent Cα atoms). In our structure, residues 523-532 at the N-terminus of the D2 domain assume an  $\alpha$ -helical conformation (equivalent to  $\alpha 2'$  of PTP1B) (46) (Figure 1A), in contrast to their disordered state in the structure of Sonnenburg et al. (35). Significantly, an equivalent  $\alpha$ -helix is present in all other cytosolic PTPs [PTP1B (46) and SHP-2 (47)] and receptor PTP structures [D1 of RPTP $\alpha$  and RPTP $\mu$ (48, 49) and D1 and D2 of LAR and CD45 (34, 50)]. In these PTP structures,  $\alpha 2'$  is preceded by helix  $\alpha 1'$ , defining the N-terminus of the tyrosine specific PTP domain (46). Multiple-sequence alignment of RPTPs (refs 33 and 34 and data not shown) predicts an equivalent  $\alpha 1'$  of RPTP $\alpha$  D2 comprising residues 502-514. These residues were not included in the RPTPa D2 domain we crystallized.

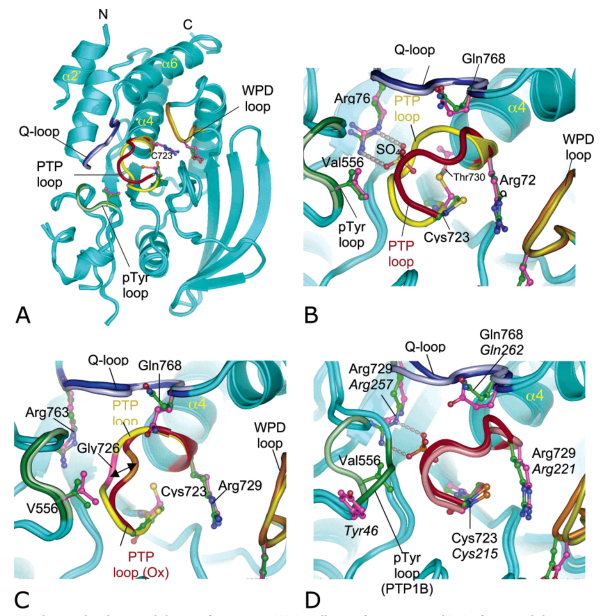


FIGURE 1: Oxidation-induced structural changes of RPTP $\alpha$  D2. (A) Overall view of RPTP $\alpha$  D2 MolA. Conformational changes are restricted to the PTP loop. Side chain carbon atoms are colored purple and green for reduced and oxidized states, respectively. Details of conformational changes of the PTP loop in MolA (B) and MolB (C). SO<sub>4</sub> represents sulfate. The change in the position of Gly726 in the PTP loop is denoted with an arrow in panel C. (D) The oxidized conformation of the PTP loop of RPTP $\alpha$  D2 MolA is identical to oxidized PTP1B (PTP1B side chains colored purple and residue numbers in italics). This figure was created using PYMOL.

Overall, the structure of RPTP $\alpha$  D2 resembles that of other tyrosine specific PTPs (Figure 1A). The domain is composed of a large central  $\beta$ -sheet flanked on both side by  $\alpha$ -helices. Residues of the PTP signature motif (Cys-Xaa<sub>5</sub>-Arg-Ser/Thr, residues 723–730) connect the central  $\beta$ -strand with the N-terminus of helix  $\alpha$ 4, creating a cradlelike structure, located at the center of the molecule, ideally configured for engaging a terminal phosphate group of a phosphorylated substrate. The side chain of the conserved Cys residue of the PTP motif (Cys723) projects into the PTP loop cradle and accepts a hydrogen bond from the side chain of Thr730 (Figure 1B).

Oxidation of RPTP $\alpha$  D2 Generates a Cyclic Sulfenamide Species. We performed a total of six oxidation experiments with RPTP $\alpha$  D2 crystals. Crystals were exposed to either 50 or 100  $\mu$ M hydrogen peroxide over a period of 16 h or oxidized glutathione for 12 days, and structures were

determined to resolutions ranging from 1.9 to 2.5 Å (Table 1). These experiments delineate the temporal oxidationdependent conformational changes of RPTPa D2. Exposure of RPTPα D2 to H<sub>2</sub>O<sub>2</sub> promotes oxidation of the conserved Cys723 residue of the PTP motif in both molecules of the asymmetric unit. This result confirms original findings that Cys723 is sensitive to oxidation (41, 42). We observe formation of the cyclic sulfenamide species in both molecules of RPTP $\alpha$  D2 on incubation with  $H_2O_2$  for 90 min (Figures 2A and 3A); however, their rates of formation and overall stability differ. Prolonged exposure (16 h) to H<sub>2</sub>O<sub>2</sub> resulted in conversion of the cyclic sulfenamide to the more oxidized SO<sub>2</sub> species in MolB, although in MolA the cyclic sulfenamide at the catalytic site remains unchanged relative to a 90 min H<sub>2</sub>O<sub>2</sub> incubation (Figures 2B and 3B). Furthermore, whereas incubation of RPTPa D2 crystals for 12 days in 10 mM oxidized glutathione generates the terminally oxidized

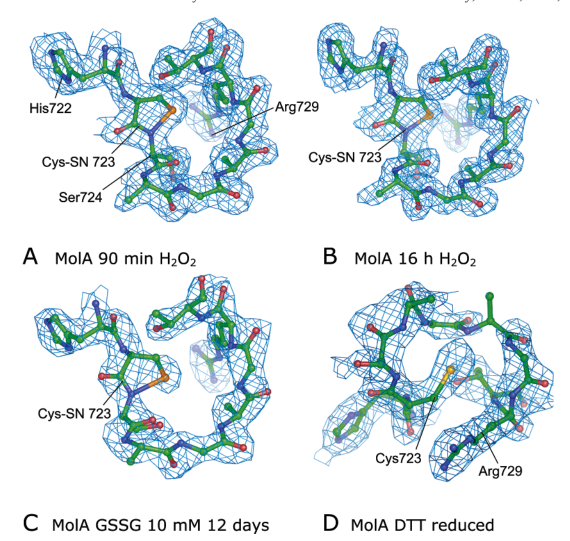


FIGURE 2: Electron density maps of oxidation states of the PTP loop of RPTPa D2 (MolA). Cyclic sulfenamide forms rapidly and is resistant to stronger oxidation. The  $2F_0 - F_c$  electron density map (contoured at  $1\sigma$ ) of the PTP loop is shown. (A) Cyclic sulfenamide forms after 90 min upon exposure to 50  $\mu$ M H<sub>2</sub>O<sub>2</sub> and is stable (B) for 16 h and (C) on exposure to 10 mM oxidized glutathione for 12 days. (D) Reduction of cyclic sulfenamide with DTT restores the native thiol. Cys-SN 723 denotes the cyclic sulfenamide species.

sulfonic acid (SO<sub>3</sub>) derivative of Cys723 in MolB (Figure 3C), a cyclic sulfenamide species is present in MolA (Figure 2C). These findings indicate significant differences in the stability of the cyclic sulfenamide between the two molecules of RPTPα D2 of the asymmetric unit. As discussed below, one explanation for this variation in stability is the difference between the PTP loop conformations in the cyclic sulfenamide states of molecules A and B. As for PTP1B (18, 19), in both reduced and oxidized states, the WPD loop adopts the open conformation.

Rates of formation of the cyclic sulfenamide species also differ between the two molecules. Following a 30 min exposure to H<sub>2</sub>O<sub>2</sub>, electron density corresponding to Cys723 is consistent with a mixture of reduced and oxidized species of the protein. In MolA, this density was modeled using partial occupancy refinement assuming a conversion of  $\sim$ 60% of molecules to the cyclic sulfenamide, with  $\sim$ 40% remaining as the reduced thiol state (Figure 1A of the Supporting Information). As in the oxidation of PTP1B, there was no evidence for a sulfenic acid intermediate, suggesting that sulfenic acid formation is rate-limiting, spontaneously condensing to the cyclic sulfenamide species. Oxidation of Cys723 in MolB is slower, with only  $\sim$ 10–20% of MolB being present as the cyclic sulfenamide at the 30 min time

point (Figure 1B of the Supporting Information). Rates of formation of the cyclic sulfenamide in RPTPα are generally similar to those in PTP1B, where complete conversion of the catalytic site Cys residue occurred within 2 h (18).

Oxidation of RPTPa D2 Is Associated with Conformational Changes. In the five-atom heterocyclic ring of the sulfenamide species, the S $\gamma$  atom of Cys723 forms a covalent bond to the main chain nitrogen atom of Ser724 and the distance between these two atoms refines to 1.5 Å (Figures 1B and 2A). The conformation of the heterocyclic ring of RPTPα D2 MolA and MolB is identical to the equivalent cyclic sulfenamide of PTP1B (18, 19) (Figure 1D). In PTP1B, formation of the cyclic sulfenamide is accompanied by profound conformational changes in the PTP loop and adjacent pTyr recognition loop. The PTP loop converts from a closed helical conformation to an open reverse  $\beta$ -hairpin that exposes the  $S\gamma$  atom of the cyclic sulfenamide species. Conformational changes of the PTP loop disrupt a hydrogen bond to Tyr46 of the pTyr recognition loop, promoting a displacement of Tyr46 to a solvent-exposed position, accompanied by shifts of the pTyr recognition loop (Figure

In RPTPα D2, the extent of conformational changes produced by oxidation to the cyclic sulfenamide species is

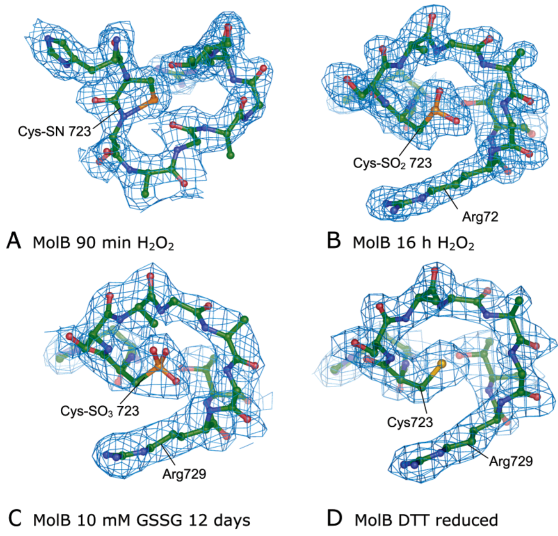


FIGURE 3: Electron density maps of oxidation states of the PTP loop of RPTP $\alpha$  D2 (MolB). The  $2F_o-F_c$  electron density map (contoured at  $1\sigma$ ) of the PTP loop is shown. Cyclic sulfenamide forms after 90 min upon exposure to 50  $\mu$ M H<sub>2</sub>O<sub>2</sub>; (B) by 16 h, cyclic sulfenamide has oxidized to SO<sub>2</sub> and (C) to SO<sub>3</sub> upon exposure to 10 mM oxidized glutathione for 12 days. (D) As in MolA, reduction of cyclic sulfenamide with DTT restores native thiol.

less pronounced than for PTP1B. In both molecules, conformational changes are confined to the PTP loop (Figure 1; Figure 2A of the Supporting Information). In MolA, the PTP loop adopts an open conformation, essentially identical to that in PTP1B, exposing the S $\gamma$  atom of the Cys723 cyclic sulfenamide (Figure 1B,D). In RPTPα D2, the PTP and pTyr loops are not connected via a hydrogen bond, and as a consequence, in contrast to PTP1B, motion of the PTP loop is not accompanied by pTyr loop conformational changes. In native, reduced RPTPα D2, the guanidinium group of a buried and invariant Arg residue of the Q-loop (Arg763, equivalent to Arg257 of PTP1B) stabilizes the closed PTP loop via hydrogen bonds with main chain carbonyl groups of PTP loop residues Ala725 and Gly726. In MolA, opening of the PTP loop disrupts these hydrogen bonds and exposes Arg763 to solvent, creating a pocket between Arg763 and the PTP loop. In the crystal, a sulfate anion engages this pocket, accepting hydrogen bonds from the side chain of Arg763 and the main chain amide of Gly726 (Figure 1B).

The oxidized PTP loop of MolB is more similar to the closed reduced conformation than to the open oxidized states of PTP1B and MolA of RPTP $\alpha$  D2. In MolB, only residue

Gly726 changes conformation by (3.6 Å) upon oxidation (Figure 1C; Figure 2A,B of the Supporting Information).

The Cyclic Sulfenamide Species Is Reduced by Thiols. As observed for PTP1B, incubation of oxidized RPTP $\alpha$  D2 crystals with 10 mM DTT for 3 h reduced the cyclic sulfenamide species to the free thiol and restored the conformation of the PTP loop to the native state (Figures 2D and 3D), demonstrating that the cyclic sulfenamide is a reversibly oxidized form of RPTP $\alpha$  D2.

The Cyclic Sulfenamide of RPTPα D2 Is Stable in Solution. Our results revealed differences in oxidation kinetics and conformational changes of the PTP loop between the two RPTPα D2 molecules of the asymmetric unit. Formation of the cyclic sulfenamide occurs earlier in MolA (within 30 min) and is resistant to stronger oxidation for at least 12 days, the maximum time of our incubation (Table 1). In contrast, in MolB, although the cyclic sulfenamide is fully formed after 90 min and remains for 6 h, as for MolA (Figure 1C,D of the Supporting Information), overnight oxidation leads to formation of sulfinic acid that is further oxidized to the sulfonic acid after 12 days (Figure 3B,C). In addition, compared with that in MolA, formation of the cyclic

sulfenamide in MolB is not associated with large conformational changes in the PTP loop (Figure 1C; Figure 2B of the Supporting Information). Differences in the kinetics of cyclic sulfenamide formation and oxidation, and conformational flexibility of the PTP loop, between MolA and MolB presumably reflect differences in their crystallographic environments.

To investigate the stability of the reversibly oxidized state of RPTPa D2 in solution over prolonged exposures to hydrogen peroxide and therefore to determine which molecule of RPTPα D2 of the asymmetric unit correlates better with the stability of the reversibly oxidized form of the phosphatase in solution, we monitored the oxidation of RPTPα D2 by means of the oxPTP antibody that recognizes the triply oxidized (SO<sub>3</sub>) state of the catalytic Cys residue (41). In the first experiment, we monitored the formation of the reversibly oxidized state in 0.1 mM H<sub>2</sub>O<sub>2</sub> over a 24 h time period using the procedure of Persson et al. (41) (sequential steps of oxidation, alkylation, reduction, and pervanadate oxidation) [Figure 4B, scheme I (ARP)]. Oxidation is maximal within 2 h of treatment and remains constant for 24 h (Figure 4A, left panel). We routinely observe a doublet upon processing of RPTPα D2, even in the RPTPα D2 C723S mutant (data not shown), indicating that doublet formation is not linked to the catalytic Cys723. However, this experiment does not distinguish between reversibly, doubly, and triply oxidized forms of the protein since all oxidized states of the catalytic Cys will convert to SO<sub>3</sub> on pervanadate incubation [Figure 4B, scheme I (ARP)]. To investigate the amount of triply oxidized RPTPa D2 (Cys-SO<sub>3</sub>) following H<sub>2</sub>O<sub>2</sub> incubation, the oxidized protein was directly blotted by the oxPTP antibody, omitting the alkylation, reduction, and pervanadate oxidation treatments. Figure 4A (right panel) indicates that very little RPTPα D2 is oxidized to the triply oxidized state even after being exposed to 0.1 mM H<sub>2</sub>O<sub>2</sub> for 24 h. To test for doubly oxidized RPTP\alpha D2 (Cys-SO<sub>2</sub>), which is not directly recognized by the oxPTP antibody, a variation of the method of Persson et al. (41) was employed [Figure 4B, scheme II (RAP)]. In this experiment, the protein was first treated with 1 mM H<sub>2</sub>O<sub>2</sub> for 30 min and then incubated with DTT to reduce reversibly oxidized cysteines (SOH, disulfides, and sulfenamides) to free thiols. Next, free thiols were alkylated, and sulfinic acid-modified cysteines were oxidized to sulfonic acid using pervanadate. The triply oxidized cysteines were then detected using the oxPTP antibody. Only very low levels of doubly oxidized RPTPα D2 are formed (Figure 5A, RAP lane), since the amount of triply oxidized RPTPa D2 following this procedure is only slightly greater than that produced following exposure to H<sub>2</sub>O<sub>2</sub> alone (Figure 5A, NP lane), consistent with MALDI-ToF data we obtained previously (42). Finally, we investigated reversible, double, and triple oxidation of RPTPα D2 in response to 0.1 mM H<sub>2</sub>O<sub>2</sub> over time and found only trace amounts of double and triple oxidation of RPTPa D2 (Figure 5B).

The time-dependent generation of oxidized species of RPTPα D2 in the presence of H<sub>2</sub>O<sub>2</sub> was also monitored using MALDI-ToF mass spectrometry (Figure 6). In this experiment, the protein was incubated in H<sub>2</sub>O<sub>2</sub>, alkylated, and then reduced (Figure 4B, scheme III). Following this procedure, reversibly oxidized cysteines will exist as reduced cysteines, whereas nonoxidized cysteines will be alkylated. As a

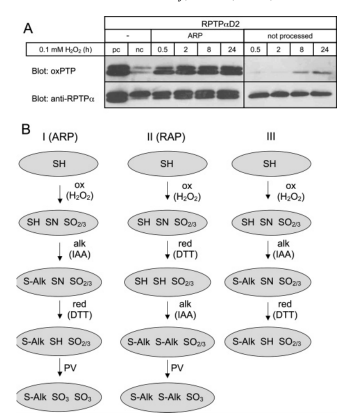
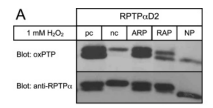


FIGURE 4: Stable reversible oxidation of RPTPα D2. (A) RPTPα D2 was treated with 0.1 mM H<sub>2</sub>O<sub>2</sub> for different periods of time as indicated and processed for detection of reversible oxidation using the oxPTP antibody as described in Experimental Procedures. The samples were run on a SDS-PAGE gel and blotted. The immunoblots were probed with the oxPTP antibody (top) and subsequently with the anti-RPTPa antibody to monitor equal loading of protein. The immunoblots were developed with enhanced chemiluminescence (ECL). The samples were processed for detection using the oxPTP antibody by alkylation, reduction, and pervanadate treatment for detection of reversible oxidation (ARP), or the samples were loaded directly on a SDS-PAGE gel for detection of irreversible triple oxidation (not processed). For the positive control (pc), the sample is treated as described for panel B [scheme I (ARP)] without steps 1 (ox) and 2 (alk). The sample is treated with DTT (red) and subsequently with pervanadate (PV), leading to maximal sulfonic acid formation. For the negative control (nc), the sample is treated as described for panel B [scheme I (ARP)], but instead of H<sub>2</sub>O<sub>2</sub> treatment in the first step, the sample is treated with 10 mM DTT for 20 min and subsequently processed (alk, red, PV), leading to maximal alkylation and minimal sulfonic acid formation. (B) Schematic of oxidation, reduction, and alkylation schemes for detection of reversibly oxidized species of RPTPa D2 in solution using the oxPTP antibody and mass spectrometry mentioned in the text and Figures 4A and 5.

positive control, RPTPα D2 was treated with pervanadate, resulting in double and triple oxidation. Figure 6 shows that during the 24 h time course of H<sub>2</sub>O<sub>2</sub> treatment the proportion of reduced to alkylated cysteine increases, whereas the amount of sulfinic and sulfonic acid remains very small, demonstrating that time-dependent reversible oxidation of Cys723 is stable.

Catalytic Activity Measurements. RPTPa D2 has some 50fold lower catalytic activity than the D1 domain, with a low affinity for pNPP ( $K_{\rm m}=21$  mM) (38). However, by measuring the catalytic activity of RPTPα D2 using 50 mM pNPP, we determined the proportion of the protein that was reversibly oxidized upon prolonged exposure to 50  $\mu$ M H<sub>2</sub>O<sub>2</sub>. Incubation of RPTPα D2 with H<sub>2</sub>O<sub>2</sub> for 18 h at 14 °C



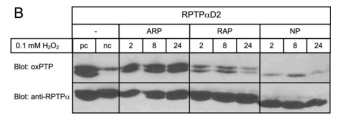
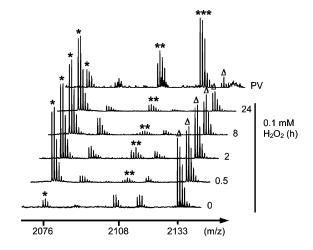


FIGURE 5: Irreversible oxidation of RPTP $\alpha$  D2 is limited. (A) RPTP $\alpha$  D2 was treated with 1 mM  $H_2O_2$  for 0.5 h and processed for detection of reversible oxidation: alkylation, reduction, and pervanadate treatment (ARP), processed for detection of irreversible oxidation, or reduction, alkylation, and pervanadate treatment (RAP), or not processed for detection of direct triple oxidation (NP). (B) Like panel A except 0.1 mM  $H_2O_2$  was used for prolonged periods of time as indicated. ARP and RAP defined in the text and the legend of Figure 4B.



- \*: singly oxidized
- \*\*: doubly oxidized
- \*\*\*: triply oxidized
  - ∆: alkylated

FIGURE 6: Reversibly oxidized RPTPa D2 is stable. Purified RPTPα D2 was treated with 0.1 mM H<sub>2</sub>O<sub>2</sub> for different periods of time (from 0 to 24 h) or as a control for irreversible oxidation with 1 mM pervanadate (PV) for 30 min. To protect the samples after H<sub>2</sub>O<sub>2</sub> and PV treatment against further oxidation during the protease digestions, the reduced cysteines were alkylated using iodoacetamide and the reversibly oxidized cysteines were reduced using DTT. Subsequently, the samples were kept under reducing conditions. MALDI-ToF spectra were acquired as described in Experimental Procedures. The m/z range that contains the active site Cys723 peptide (residues 710-729) is depicted. The reversibly oxidized Cys723 peptide was detected in reduced form (m/z 2075.98, one asterisk); the sulfinic acid-containing peptide shifted +32.00 to m/z2107.98 (two asterisks), and the sulfonic acid-containing peptide shifted +48.00 to m/z 2123.98 (three asterisks). The reduced Cys723-containing peptide was detected in alkylated form with a shift in relative molecular mass of +57.02 (m/z 2133.00,  $\Delta$ ).

completely inactivated the enzyme. Exposure of the oxidized protein to 25 mM DTT for 3 h restored the specific activity to 72.5% (mean of two experiments, range from 72.4 to 72.5%) of its pre-oxidation activity. In control experiments

in the presence of 5 mM DTT, no loss of phosphatase activity was observed after 21 h. These results indicate that the majority of RPTP $\alpha$  D2 exists as a reversibly oxidized state on prolonged exposure to  $H_2O_2$ . Failure to recover all catalytic activity on reduction with DTT suggested that a small proportion of the protein was irreversibly oxidized, most likely modified to the higher oxidized sulfinic and sulfonic states, in agreement with the Western blot studies showing small amounts of these species on exposure to 100  $\mu$ M  $H_2O_2$  for 24 h (Figures 4A and 5B).

Together, these results demonstrate that in solution, the majority of RPTP $\alpha$  D2 ( $\sim$ 72%) exists in the reversibly oxidized state on prolonged exposure to H<sub>2</sub>O<sub>2</sub> (18–24 h) and is therefore resistant to stronger oxidation. This correlates better with the stability of the cyclic sulfenamide observed in MolA of the crystal structure. Whereas in MolA, electron density consistent with the cyclic sulfenamide is observed on 16 h exposure to H<sub>2</sub>O<sub>2</sub>, in MolB Cys723 was oxidized to sulfinic acid (Figures 2B and 3B).

RPTPα D2 Is a Monomer in Solution. Using calibrated gel filtration chromatography, we determined the apparent molecular mass of RPTPα D2 to be 28 kDa (calculated mass of 31 kDa) (data not shown), consistent with a monomeric state of the protein in solution. Our findings confirm those of Sonnenburg et al., who determined that RPTPa D2, as well as the RPTPα D1D2 protein, is a monomer in solution as assessed by gel filtration and dynamic light scattering (35). The apparent molecular mass of RPTPα D2 did not change on conversion to the oxidized state, indicating that oxidation of the phosphatase did not promote protein dimerization. The dimers of RPTPa D2 we observe in the crystal, similar to that of Sonnenburg et al. (35) and favored by the crystallization conditions, may contribute to the interaction interface within constitutive RPTPa dimers observed within a cellular context (51, 52). However, additional factors, such as the involvement of other domains of RPTPa and/or the membrane environment, are likely to stabilize RPTPα dimers in vivo.

#### **DISCUSSION**

The Cyclic Sulfenamide as a Universal Feature of Reversibly Oxidized Conventional PTPs. In this study, we demonstrate that RPTPa D2 is reversibly oxidized via a cyclic sulfenamide species. The cyclic sulfenamide is promoted by oxidants H<sub>2</sub>O<sub>2</sub> and oxidized glutathione and reduced by DTT. The discovery of the cyclic sulfenamide in RPTPα D2 is only the second example of such a species in any protein and the first report of a RPTP D2. However, the existence of a cyclic sulfenamide in two distinct PTP domains suggests that its formation is not unique to PTP1B (18, 19) but will likely be a general feature of catalytic Cys oxidation in conventional PTPs. Oxidation of DSPs, lmPTPs, Cdc25, and lipid phosphatases that feature cysteine residues in the proximity of the catalytic Cys promotes disulfide bond formation (5). Interestingly, only one conventional PTP, mPEP, shares a proximal Cys equivalent to those of KAP and PTEN (33), and it will be interesting to determine whether a cyclic sulfenamide or disulfide bond represents the reversibly oxidized state of this protein.

The role of the catalytic Cys of RPTP D2s as a redox sensor may explain their conservation in domains with little or no phosphatase catalytic activity (4). Structural features of the phosphatase catalytic site are conserved in D2s to create the microenvironment necessary for the reduced p $K_a$ of the catalytic Cys residue essential for its role as a redox sensor. These include a positive electrostatic potential created by the conserved Arg residues of the PTP motif and the Q-loop (Arg729 and Arg763, respectively), the hydrogen bond between side chains of Cys723 and Thr730 of the PTP loop, and the orientation of the main chain amides of the PTP loop toward the thiol of Cys723 (Figure 1B). Kinetic data indicate that the Arg residue of the PTP C(X)<sub>5</sub>R motif together with a histidine residue immediately N-terminal to the PTP motif, conserved in PTPs, contributes to the reduced  $pK_a$  of the catalytic Cys residue (16, 53). Interestingly, the PTP signature motif of conventional tyrosine specific PTPs comprises three highly conserved glycine residues: Cys-Ser-Xaa-Gly-Xaa-Gly-Arg-(Ser/Thr)-Gly. The conformational flexibility conferred by glycines may permit opening of the PTP loop and stabilization of the cyclic sulfenamide. Both the PTP loops of PTP1B and RPTP $\alpha$  D2 MolA adopt the same open conformation upon formation of the cyclic sulfenamide, and in RPTP $\alpha$  D2, this is associated with its resistance to stronger oxidation observed in the crystal.

Implications for Oxidation-Dependent RPTPa Conformational Changes and Regulation. A mechanism for regulating RPTP catalytic activity involving formation of inhibitory RPTP dimers, proposed on the basis of the crystal structure of RPTPa D1 dimers, in which wedge elements of each subunit insert into the catalytic site of the opposing monomer (48), is consistent with a range of biochemical and cellular studies (54-56). Significantly, the structure explained earlier observations that signaling from a chimera of the extracellular and transmembrane regions of EGFR fused to the cytosolic domain of CD45 was negatively regulated by EGF-induced dimerization (54). However, crystal structures of the LAR and CD45 tandem D1 and D2 domains showed that formation of the inhibitory D1 interface would be prevented by a steric clash between the D2 domains of the dimer, which could be relieved only by conformational changes which alter the D1-D2 domain interface (34, 50). The relative orientation of the LAR and CD45 D1 and D2 domains is similar, constrained by a short four-residue linker, and stabilized by a set of extensive interactions (Figure 7A). LAR and CD45 residues that mediate interdomain contacts, together with the linker region, are highly conserved in RPTP $\alpha$  (34), suggesting that the relative orientation of D1 and D2 in RPTPα will be similar to that in LAR and CD45 (Figure 7).

Substantial evidence now indicates that oxidation of Cys723 of RPTPa causes changes in RPTPa tertiary and quaternary structures. Oxidation induces changes in FRET between GFP derivatives attached to the N-terminus of α1' and the C-terminal  $\alpha$ -helix of RPTP $\alpha$  D2 ( $\alpha$ 6, Figure 7A) (39) and alters the protein's sensitivity to trypsinolysis (A. Groen, J. Overvoorde, L. Tertoolen, T. van der Wijk, and J. den Hertog, unpublished observations). These tertiary structural changes are associated with changes in quaternary structure in such a way that preexisting RPTPa dimers are stabilized and their rotational coupling is induced (39, 40).

In RPTPα D2 crystals, oxidation-dependent conformational changes are confined to the PTP loop. Unlike that in PTP1B, the neighboring pTyr loop is not altered, and therefore, conformational changes of the PTP loop are not

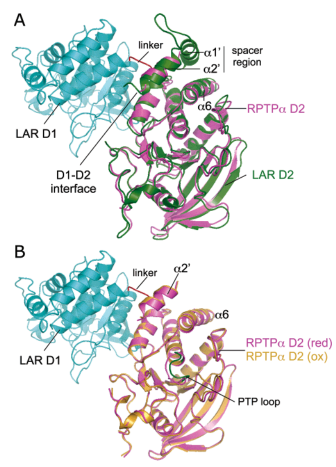


FIGURE 7: Consequences of RPTPα D2 oxidation for the D1–D2 interface. Superimposition of RPTPa D2 (reduced and oxidized) onto D2 of LAR. (A) D2 of RPTPa (purple) superimposed onto D2 (green) of LAR. The linker region of LAR is colored red, and LAR D1 is colored blue. (B) Oxidized and reduced RPTPa D2 (MolA) relative to D1 of LAR. LAR D2 has been omitted for clarity. Oxidation-induced conformational changes of the RPTP $\alpha$ D2 catalytic site do not directly alter the D1-D2 interdomain interface.

directly transmitted to residues of RPTPa D2 that, on the basis of sequence conservation with LAR and CD45, would be predicted to participate at the D1-D2 domain interface (Figure 7B). The more extensive oxidation-dependent conformational changes of RPTPα D2 required an explanation of oxidation-dependent suppression of RPTPα activity, and coupled tertiary and quaternary conformational changes that are not observed in our crystallographic study. One possible explanation is that crystal lattice contacts limit the capacity of RPTPa D2 to undergo conformational change in response to oxidation. Our finding that the two molecules of RPTPa D2 respond differently to H<sub>2</sub>O<sub>2</sub> supports the notion that lattice contacts influence the extent of conformational change. Furthermore, our efforts to crystallize oxidized RPTPα D2 were unsuccessful, a result which could be explained if the oxidized protein adopted a conformation different from those observed in the crystal. It is also possible that the absence of the putative α1' helix in our RPTPα D2 crystals could prevent large conformational changes if  $\alpha 1'$  were required to stabilize the oxidized conformation of RPTP $\alpha$  D2. Alternatively, an allosteric mechanism could be invoked to explain how oxidation of Cys723 promotes changes at the D1-D2 interface. In this model, more extensive RPTPα tertiary and quaternary conformational changes promoted by

Cys723 oxidation occur only in the context of the entire RPTPα protein. The RPTPα extracellular and transmembrane domains are independently capable of homodimerization (*51*, *52*), whereas RPTPα D2 homodimerizes upon Cys723 oxidation (*39*). Oxidation of Cys723 (presumably by causing conformational changes of the PTP loop altering the protein's molecular surface) promotes D2–D2 associations (*39*) affecting the overall quaternary structure of RPTPα. The oxidation-induced association of D2 domains could be triggered by changes in the surface properties of D2 resulting from generation of the cyclic sulfenamide and opening of the PTP loop or, as suggested by van der Wijk et al. (*57*), be mediated by intermolecular disulfides involving Cys723.

In conclusion, our data demonstrate that oxidation of Cys723 of RPTP $\alpha$  D2 yields a reversible cyclic sulfenamide species, associated with conformational changes in the PTP loop. However, these limited structural changes are insufficient for explaining  $H_2O_2$ -mediated dimerization, and more extensive tertiary conformational changes within RPTP $\alpha$  D2 may be expected in the context of the full-length RPTP $\alpha$  protein, and without the constraints imposed by the crystal lattice.

#### **ACKNOWLEDGMENT**

We thank J. P. Noel (The Salk Institute for Biological Studies, La Jolla, CA) for the RPTP $\alpha$  D2 expression plasmid. We thank staff at ESRF for help with data collection.

#### SUPPORTING INFORMATION AVAILABLE

Electron density maps of various oxidation states of the PTP loop of RPTP $\alpha$  D2 (Figure 1) and oxidation-induced structural changes in RPTP $\alpha$  D2 (Figure 2). This material is available free of charge via the Internet at http://pubs.acs.org.

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BI061546M