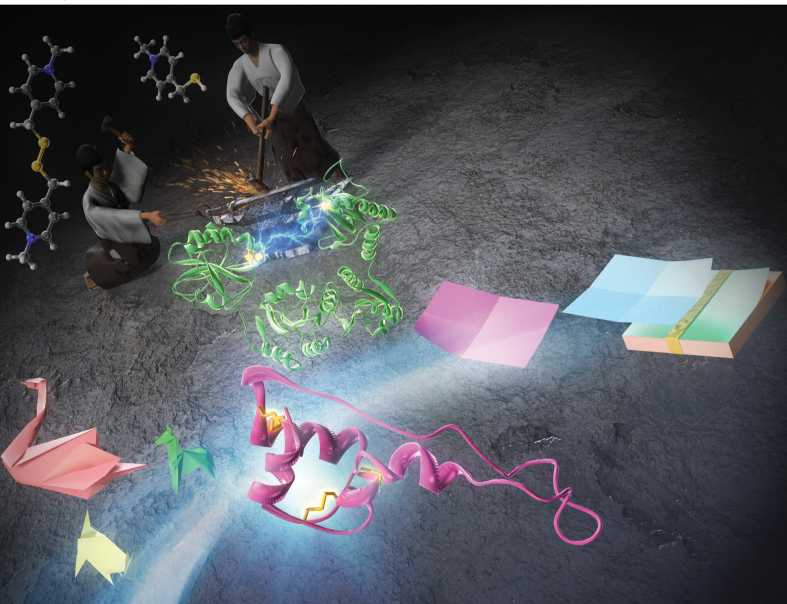
Volume 60 Number 48 18 June 2024 Pages 6073-6212

ChemComm

Chemical Communications

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ISSN 1359-7345



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Takahiro Muraoka, Masaki Okumura *et al.* Boosting the enzymatic activity of CxxC motif-containing PDI family members



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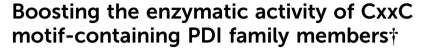


Cite this: Chem. Commun., 2024, 60, 6134

Received 11th April 2024, Accepted 9th May 2024

DOI: 10.1039/d4cc01712a

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Tsubura Kuramochi,‡^{ab} Yukino Yamashita,‡^c Kenta Arai, pde Shingo Kanemura, pda

Compounds harboring high acidity and oxidizability of thiol groups permit tuning the redox equilibrium constants of CxxC sites of members of the protein disulphide isomerase (PDI) family and thus can be used to accelerate folding processes and increase the production of native proteins by minimal loading in comparison to glutathione.

Chemical boosters of enzymatic activity provide us with a better understanding of enzymatic function and allow us to combat enzyme-related pathologies. Members of the protein disulphide isomerase (PDI) family are responsible for catalyzing oxidative protein folding in the endoplasmic reticulum (ER).¹⁻³ Several studies have reported that PDIs play important roles in protein misfoldingrelated pathologies, such as amyotrophic lateral sclerosis, Alzheimer's disease (AD), Parkinson's disease, and type-II diabetes. 4,5 Lossof-function of PDIs can thus result in severe diseases, such as neurodegeneration and diabetes. Notably, redox active site dysfunction in PDIA1 and PDIA6 CxxC motifs induced by posttranslational chemical modifications has been observed in AD,6 suggesting that misfolding-related pathologies can be ameliorated using redox chemistry.7

Extracellular PDIs are promising targets for thrombosis-related and tumor diseases owing to the upregulation of the PDI expression

level, and several PDI inhibitors, such as 16F16, bisphenol A, PACMA31, LOC14, and piericone, have been reported. 8-12 Many of these antagonists abolish the PDI catalytic activity by acting on cysteines within CxxC motifs or by competitively inhibiting its function by binding to its substrate-binding pocket. However, in the case of PDIA1, an allosteric switch of its substratebinding pocket increases its catalytic activity; 13 yet, only a few studies were reported about an enhancer of PDIs activity. 16 Regarding thiol compound-mediated catalytic activity, 17 developing novel thiol compounds to target cysteines within CxxC motifs could be extremely effective.

In the ER, glutathione has a central role in thiol-disulphide exchange reactions, where both its reduced (GSH) and its oxidized (GSSG) form catalyze oxidative protein folding¹⁸ and control the PDIs redox status.¹⁹ Regarding thiol compounds that can replace glutathione (GSH; $pK_a = 9.17$, $E^{\circ\prime} = -256$ mV), we previously reported that para-substituted N-methylated pyridinylmethanethiol (pMe-PySH; p $K_a = 7.34$, $E^{\circ\prime} = -211$ mV) enhances both the acidity and oxidizability of the thiol groups, which allows several clients to accelerate oxidative folding by 1-equivalent loading as semienzymatic activity (Fig. 1A).20 Such a highly redox-reactive function of pMePySH by minimum loading prompted us to explore how the PDIs enzymatic activity can be controlled via the CxxC motif.

Among PDIs, PDIA1, PDIA6, and PDIA15 are physiologically involved in the oxidative folding of proinsulin and insulin secretion, 21 and PDIA3 is also essential for the efficient folding of glycoproteins in cells.22 We herein validated a strategy via the CxxC motif of PDIA1, PDIA3, PDIA6, and PDIA15 due to the redox-related pathological significance. To evaluate the "redox" molecule that enhances enzymatic activity by minimal loading, we investigated its effect on the PDI active sites when its concentration was in the µM rather than in the mM range. 19 Although the thiol-disulphide catalytic CxxC motif is highly conserved in the PDI family (Fig. S1 in ESI†), the reactivity of each active site with glutathione differs depending on the structural characteristics of the PDIs (Fig. 1B, C and Fig. S2, ESI†). The redox equilibrium constant of each PDI member was determined by incubating purified PDIs in

^a Frontier Research Institute for Interdisciplinary Sciences, Tohoku University, 6-3 Aramaki-Aza-Aoba, Aoba-ku, Sendai, Miyagi, 980-8578, Japan. E-mail: okmasaki@tohoku.ac.jp

^b Department of Molecular and Chemical Life Sciences, Graduate School of Life Sciences, Tohoku University, 2-1-1 Katahira, Aoba-ku, Sendai, Miyagi, 980-8577, Japan

^c Department of Applied Chemistry, Graduate School of Engineering, Tokyo University of Agriculture and Technology, 2-24-16 Naka-cho, Koganei, Tokyo 184-8588, Japan. E-mail: muraoka@go.tuat.ac.jp

^d Department of Chemistry, School of Science, Tokai University, Kitakaname, Hiratsuka-shi, Kanagawa 259-1292, Japan

^e Institute of Advanced Biosciences, Tokai University, Kitakaname, Hiratsuka-shi, Kanagawa 259-1292, Japan

^fKanagawa Institute of Industrial Science and Technology, 3-2-1 Sakato, Takatsu-ku, Kawasaki, Kanagawa 213-0012, Japan

[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d4cc01712a

[‡] These authors contributed equally to this work.

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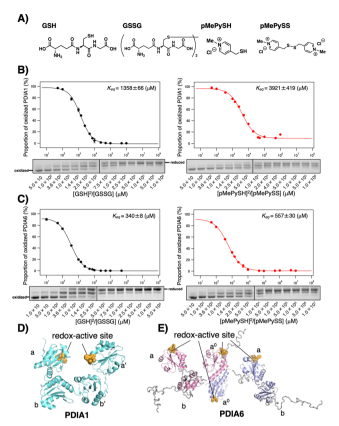


Fig. 1 Redox equilibrium constants of PDIA1 and PDIA6 using different GSH/ GSSG or pMePySH/pMePySS ratios. (A) Chemical structures of GSH, GSSG, pMePySH, and pMePySS. Redox equilibrium constants of (B) PDIA1 and (C) PDIA6 using different GSH/GSSG or pMePySH/pMePySS ratios. Free PDI thiol groups (0.3 μ M) were modified with malPEG2000 after incubation with different [GSH]²/[GSSG] or [pMePySH]²/[pMePySS] ratios in a degassed buffer containing 90 μM GSSG or pMePySS and various concentrations of GSH or pMePySH (30-30000 $\mu M)$ at 30 °C. \textit{K}_{eq} values were determined from at least three independent experiments. (D) PDIA1 consists of four Trx-like domains, the first and last of which contain redox-active CxxC motifs. PDIA1 (PDB code; 4EKZ) crystal structure revealing four domains, which form an overall U-shaped structure.¹⁴ (E) PDIA6 consists of three Trx-like domains, the first and second of which contain redox-active CxxC motifs. PDIA6 dimerizes in solution via a unique dimeric motif.¹⁵ The dimeric motif is comprised of a Leu- and Val-rich region in the first redox active a⁰ domain. Of note, in contrast to PDIA1, the solvent-exposed redox active sites in each PDIA6 Trx-like domain rapidly introduce disulphide bond(s) into the client proteins. This structural feature is quite different from that of PDIA1.

redox buffers containing different ratios of [GSH] to [GSSG] or [pMePySH] to [pMePySS]. To monitor the number of free thiol(s) of Cys residue(s), the reaction mixtures were modified by malPEG2000 (average $M_{\rm n}$ = 2000) and subsequently separated by SDS-PAGE (Fig. S2C, ESI†). The apparent redox equilibrium constants of four representative PDIs (PDIA1, PDIA3, PDIA6, and PDIA15) were calculated from the dose-response curve for redox agents obtained from the stained gel intensity (Fig. 1B, C and Fig. S3, ESI†). Consistent with a previous report, 19 PDIA1 exhibited the highest redox equilibrium constant (K_{eq} = 1358 μ M), whereas PDIA6 exhibited the lowest constant ($K_{eq} = 340 \mu M$). The PDIA1 redox active sites face the interior of the U-shaped structure, 1,14 whereas the same sites in PDIA6 are exposed to the solvent, 15 resulting in significantly different reactivity between the PDIs and glutathione active sites (Fig. 1D and E). As for pMePySH/pMePySS, the ranking of the redox equilibrium constants among PDIs was the same as those of GSH/GSSG (Fig. 2). However, among four representative PDI family members, the redox equilibrium constants using pMePySH/pMePySS were higher than those using GSH/ GSSG, indicating higher reactivity with the redox active CxxC sites within the PDI family (Fig. 2).

To validate the ability to catalyze the oxidative folding against a reduced and denatured client, the oxidative folding of bovine pancreatic trypsin inhibitor (BPTI) was monitored in the presence of disulphide/thiol compounds ([BPTI] = 30 μM; [disulphides] = 90 μ M; [thiols] = 360 μ M). Native BPTI has three pairs of cysteine residues that can form disulphide bonds, i.e., Cys5-Cys55, Cys14-Cys38, and Cys30-Cys51 (Fig. 3A); therefore, since pMePySS accelerates the oxidative folding most efficiently by 1-equivalent loading as semi-enzymatic activity, 20 we used a 90 µM disulphide concentration. Considering that the optimal condition for PDIA1 activity is [GSH]: $[GSSG] = 5:1^{17}$ and that a [GSH]:[GSSG] = 3:1 may be present in the ER, 23 [thiols] = 360 μ M and [disulphides] = 90 μ M were used. Regarding on-path BPTI folding (Fig. 3B), quasi-native intermediates such as N' and N^* fold into native-like structures with two disulphide bonds and form native structures (N) via $N_{\rm SH}^{\rm SH}$. In the absence of PDIs, BPTI folds relatively slowly, with yields of 34.7% for GSH/GSSG and 72.6% for pMePySH/pMe-PySS in 120 min (Fig. S4, ESI†). Consistent with previous reports, 1,2 BPTI folding was accelerated by PDIA1 or PDIA6 under GSH/GSSG, with yields of 90.4% for PDIA1 and 58.7% for PDIA6 in 120 min (Fig. 3C, F and Table 1). In comparison to GSH/GSSG, pMePySH/pMePySS enhanced the BPTI folding efficiency, which resulted in an N yield of 96.5% for PDIA1 and 80.7% for PDIA6 after 120 min (Fig. 3D and G). To further calculate the BPTI folding kinetics, the rate constants for BPTI folding were determined to be $k_{\text{GSH-PDIA1}} = 2.3 \times 10^{-2} \text{ min}^{-1}$, $k_{\text{pMePySH-PDIA1}} = 9.0 \times 10^{-2} \text{ min}^{-1}, k_{\text{GSH-PDIA6}} = 1.6 \times 10^{-2} \text{ min}^{-1},$ and $k_{\text{pMePySH-PDIA6}} = 4.1 \times 10^{-2} \text{ min}^{-1}$, *i.e.*, the PDI-catalyzed reaction rate using pMePySH/pMePySS was indeed 2.5-4.0 times faster than that using the typical GSH/GSSG system (Fig. 3E, H and Fig. S5, ESI†). Of note, PDIA1 and PDIA6 significantly

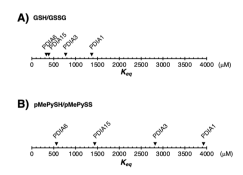


Fig. 2 Redox equilibrium constant ($K_{\rm eq}$) of PDI family members using different (A) GSH/GSSG or (B) pMePySH/pMePySS ratios. The K_{eq} values of PDIA1, PDIA3, PDIA6, and PDIA15 using pMePySH/pMePySS were determined as 3921, 2809, 1441, and 557 μ M, respectively.

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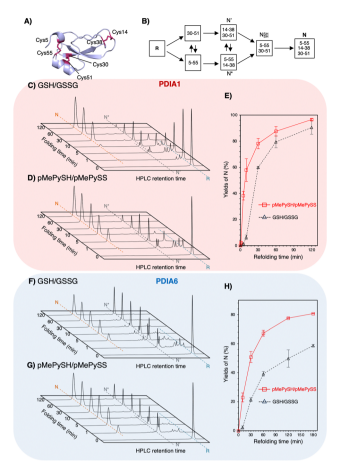


Fig. 3 Both PDIA1 and PDIA6 under the pMePySH/pMePySS redox system accelerated the oxidative folding rates. (A) The crystal structure of BPTI (PDB code; 6PTI). The natively folded BPTI has three disulphide bonds (Cys5-Cys55, Cys14-Cys38, and Cys30-Cys51). (B) Oxidative folding pathway of BPTI. Time-course reverse-phase HPLC analyses of oxidative folding of BPTI (30 μ M) in the presence of: (C) PDIA1 (0.3 μ M) under a GSH (360 μ M)/GSSG (90 μ M) redox system, (D) PDIA1 (0.3 μ M) the pMePySH (360 μ M)/pMePySS (90 μ M) redox system, (F) PDIA6 (0.3 μ M) under a GSH (360 $\mu\text{M})/\text{GSSG}$ (90 $\mu\text{M}) redox system, and (G) PDIA6 (0.3 <math display="inline">\mu\text{M})$ under the pMePySH (360 μ M)/pMePySS (90 μ M) redox system, respectively. (E) and (H) Time-course plots of native BPTI yields in (C), (D), (F) and (G). N, N', N*, and R represent native, folding intermediates with 2-disulfide pairings $(30-51 \text{ and } 5-55 \text{ for } N', \text{ and } 5-55 \text{ and } 14-38 \text{ for } N^*), \text{ and reduced forms}$ of BPTI. Eluent buffers, water (containing 0.1% TFA) and CH₃CN (containing 0.1% TFA) with a linear gradient; flow rate, 1.0 mL min⁻¹; detection wavelength, 229 nm; temperature, 30 °C.

accelerated the production of N in BPTI under the redox environment using pMePySH/pMePySS (Fig. 3).

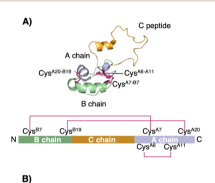
To further ensure the generality of the substrate catalyst, we investigated its ability to promote oxidative folding of human proinsulin, a peptide of biopharmaceutical interest for diabetes treatment. Human proinsulin consists of three chains, A, B, and the so-called C peptide, which connects A and B, with two disulphide bonds (Cys^{B7} – Cys^{A7} and Cys^{B19} – Cys^{A20}) residing between chains A and B and one (Cys^{A6} – Cys^{A11}) constituting an intra-A-chain bond (Fig. 4A). The three disulphide bonds are crucial for the native folding of mature insulin, but we observed that its refolding yield in the μM range of GSH/GSSG was grossly impaired at neutral pH (Fig. 4B). Therefore, PDIs are

Table 1 BPTI native-form yields after 120 min refolding time

Combination of enzymes and redox agents	Native BPTI yields ^a (%)
GSH/GSSG	34.7 ± 1.0
pMePySH/pMePySS	72.6 ± 4.3
PDIA1+GSH/GSSG	90.4 ± 5.1
PDIA1+pMePySH/pMePySS	96.5 ± 0.8
PDIA6+GSH/GSSG	58.7 ± 0.9
PDIA6+pMePySH/pMePySS	80.7 ± 0.2

 $[^]a$ Yields were calculated from the area of RP-HPLC. Errors indicate the mean \pm SEM of three independent experiments.

thought to be involved in proinsulin oxidative folding under physiological conditions, ²¹ and PDI-mediated catalytic activity enhancement is extremely important for insulin production. Compared with the GSH/GSSG system, both PDIA1 and PDIA6 with pMePySH/pMePySS increased the efficiency of native formation of human proinsulin by approximately 10%. Remarkably, GSH/GSSG



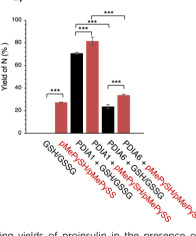


Fig. 4 Refolding yields of proinsulin in the presence of GSH/GSSG or pMePySH/pMePySS with/without PDIA1 or PDIA6. (A) Solution structure of proinsulin (PDB code; 2KQP). The natively folded proinsulin has three disulphide bonds (Cys $^{\rm B7}$ –Cys $^{\rm A7}$, Cys $^{\rm B19}$ –Cys $^{\rm A20}$ and Cys $^{\rm A6}$ –Cys $^{\rm A11}$). (B) The yields of native proinsulin (5 μ M) in the presence of GSH/GSSG or pMePySH/pMePySS with/without PDIA1 (0.5 μ M) or PDIA6 (0.5 μ M) after 5 min are shown. Oxidative folding was performed at pH 7.5. Before HPLC analyses, aliquots of the folding solutions were pre-treated with aqueous 2-aminoethyl methanethiosulfonate (AEMTS) to quench the reaction by blocking the free SH groups into –SSCH₂CH₂NH₃+. Error bars indicate the mean \pm SEM of three independent experiments. The statistical significance of differences was examined by one-way analysis of variance with Tukey honestly significant difference (HSD) post hoc testing. All statistical tests were performed using KaleidaGraph statistical software (Synergy Software) at a significance level of α = 0.05.

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alone failed to increase the native form of proinsulin, but PDIA1 in the µM range of pMePySH/pMePySS increased the yield by up to 81.6% (Fig. 4B). Thus, PDIA1 and PDIA6 markedly increased the yield of the protein native structures by minimal loading of pMePySH in comparison to glutathione.

Differences in redox equilibrium constants among individual PDI family members are believed to sustain the complicated but elaborate redox network in the ER that controls protein homeostasis. 19 Therefore, pathological redox imbalances can be caused by uncontrolled oxidation (e.g., excessive amount of reactive oxygen species) or reduction (e.g., hypo-oxidation in the ER contributing to the etiology of misfolding diseases such as diabetes and AD24). Further, dysfunction of PDI family members is known to cause pathological diseases such as neurodegeneration and diabetes.^{4,7} Therefore, chemically controlling the PDI family as ER-resident redox enzymes is crucial.

Here, we found that pMePySH/pMePySS redox systemmediated catalysis accelerated the folding kinetics of PDIA1 (with a relatively high K_{eq}) and PDIA6 (with a relatively low K_{eq}). This work could offer a proof of concept regarding the existence of redox crosstalk in the ER, i.e., thiol compounds targeting the cysteines of PDI family CxxC motifs and increasing the PDI family member enzymatic activity more efficiently than glutathione, a typical redox agent of universal use. Regarding the chemical booster of PDI activity, an allosteric switch operating via the substrate-binding b' domain increases PDI activity, 13 but a similar allosteric effect can also lead to decreased activity.11 One reason for this apparent paradox is that regulation of the both function and structure is beyond the proof-ofconcept for chemical design due to the conformational dynamic nature of the PDI family. 1,25 Therefore, chemical boosters of the PDI family activity via CxxC motif(s) may further provide clues to new approaches for combating PDI familyrelated misfolding pathologies.

MO conceived the idea of this study. TK, YY, SK, and KA performed the experiments. MO and TM supervised the implementation of this study. MO wrote the manuscript. All authors critically reviewed and revised the manuscript draft and approved the final version for submission.

This research was funded by a JSPS KAKENHI grant (grant numbers 22H02205 (to MO)), a Fund for the Promotion of Joint International Research (grant number 23KK0105 (to MO)), JSPS Grants-in-Aid for Transformative Research Areas (B) (grant numbers JP21H05095 (to MO) and JP21H05096 (TM)), the Japan Science and Technology Agency FOREST Program (grant numbers JPMJFR201F (to MO) and JPMJFR2122 (to TM)), a grant from the Takeda Science Foundation (to MO and TM), a grant from the Mochida Memorial Foundation for Medical and Pharmaceutical Research (to MO), a grant from the Naito Foundation (to MO), a grant from the Uehara Memorial Foundation (to MO), a grant from the Terumo Life Science Foundation (to MO), a grant from the Astellas Foundation for Research on Metabolic Disorders (to MO), a grant from The Asahi Glass Foundation (to MO and TM), Mitsui Sumitomo Insurance Welfare Foundation (to MO), a grant from the Daiichi Sankyo Foundation of Life Science (to MO), and a grant from THE Sumitomo FOUNDATION (to MO). This work was supported by

JST SPRING, Grant number JPMJSP2114. Generous support was received from the FRIS CoRE, which is a shared research environment at Tohoku University. We thank Prof. Kenji Inaba (Kyushu University) for the kind gift of PDI family plasmids.

Conflicts of interest

There are no conflicts of interest to declare.

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