Chapter 9

Analysis of the Mechanism That Brings Protein Disulfide Isomerase-P5 to Inhibit Oxidative Refolding of Lysozyme

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Abstract Mammalian sperm acquire fertilizing capacity during epididymal transit. During this process, the expression of protein disulfide isomerase-P5 (PDI-P5, P5) decreases. Almost all members of the PDI family accelerate the formation, reduction, and isomerization of disulfide bonds, whereas boar P5, which is composed of two active thioredoxin domains, \(a\) and \(a'\), and an inactive \(b\) domain, inhibits the oxidative refolding of reduced and denatured lysozyme due to nonproductively folded lysozyme resulting from intermolecular isomerization. We investigated the reductive activities of cysteine mutants of boar P5 and of the \(a\) and \(a'b\) domains, using an insulin turbidity assay, and their ability to inhibit the oxidative refolding of reduced and denatured lysozyme. We also analyzed refolded products by Western blotting. The reductive activities of the C-terminal variants, C171A, C174A, and C171/174A, were lower than those of the N-terminal variants C36A, C39A, C36/39A, and \(a'b\). The inhibitory activities of N- and C-terminal cysteine mutants were \(~60\%\) and \(~75\%\), respectively, whereas the inhibitory activities of C36/39A and of \(a'b\) were \(~60\%\) and \(30\%\), respectively. The properties of a mixture of \(a\) and \(a'b\) were similar to those of \(a'b\) alone. These results suggested that two active domains of P5 cooperatively inhibited refolding by intermolecular isomerization and that this effect was increased when the \(a\) and \(a'b\) domains were linked. The findings were supported by Western blotting analysis of the refolded products. These results provide new insight into the molecular mechanism by which P5 inhibits protein refolding.
9.1 Introduction

Caput epididymal sperm released from the testes are biologically incompetent, acquiring functionality during transit from the epididymal caput to the cauda via the corpus. Ejaculated sperm are able to move vigorously and participate in the complex cascades of interactions that culminate in fertilization of the oocyte (Yanagimachi et al. 1994). Various kinds of studies on epididymal sperm maturation have been reported (McLaughlin et al. 1997; Frayne et al. 1998; Zhu et al. 2001; Saxena et al. 2002; Baker et al. 2005, 2012; Ellerman et al. 2006; Toshimori et al. 2006; Inoue et al. 2008; Ijiri et al. 2011). Protein disulfide isomerase-P5 (PDI-P5, P5) has been reported present in mouse sperm membrane fraction (Stein et al. 2006) and to be involved in asymmetrical organogenesis in zebrafish (Hoshijima et al. 2002), in regulating human platelet function (Jordan et al. 2005), and in promoting the ability of human tumors to evade the immune system (Kaiser et al. 2007; Gumireddy et al. 2007). We have reported that boar P5 is downregulated during transit from the epididymal corpus to the cauda and inhibits oxidative refolding of reduced and denatured lysozyme (Akama et al. 2010). To date, however, the mechanism by which P5 inhibits protein refolding has not been elucidated.

P5 is composed of two active thioredoxin domains, $\alpha$ and $\alpha'$, and an inactive $\beta$ domain, with each active thioredoxin domain containing an active CGHC site. We investigated the mechanism by which P5 inhibits protein refolding by assaying the inhibitory activities of highly purified domain-deleted or cysteine variants of P5. Our results suggest that the $\alpha$ domain inhibits refolding by promoting the intermolecular isomerization of lysozyme with the $\alpha'$ domain, with the $\alpha$-$\alpha'$-$\beta$ attachment essential for complete inhibition.

9.2 Materials and Methods

9.2.1 Expression and Purification of PDI-P5 Variants

The cDNAs corresponding to P5 deletion variants, the $\alpha$ and $\alpha'b$ fragments, were amplified by polymerase chain reaction (PCR) and inserted into the expression vector pET30a(+) (Novagen). The cysteine mutants of P5 cDNA were generated by amplification of wild-type P5 cDNA cloned into pET30a(+) vector using pfu polymerase (Promega). The domain structure of P5 and its variants are shown in Fig. 9.1. Expression and purification of pET-PDI-P5 variants were performed as described (Akama et al. 2010), with ammonium sulfate precipitation and hydrophobic chromatography on phenyl-Sepharose (CL-4B) (GE Healthcare Bio-Science) performed before affinity chromatography.
9.2.2 Insulin Turbidity and Lysozyme Refolding Assays

Insulin turbidity (Holmgren 1979) and lysozyme refolding (Puig and Gilbert 1994) assays were performed as described.

9.2.3 Western Blotting

Reaction products of lysozyme refolding assay were Western blotted as described (Akama et al. 2010).

9.3 Results

9.3.1 Reductive Activity of $a'$ Domain

Purified recombinant mature P5 showed thiol-dependent reductase activity, catalyzing the reduction of insulin disulfides with DTT as previously reported (Akama et al. 2010). All cysteine-to-alanine mutations decreased the reductive activity of P5.
C-terminal mutants (C171A, C174A, and C171/174A) had about ~50 % of the activity of wild type, whereas N-terminal mutants (C36A, C39A, C36/39A and \(a'b\)) had about ~75 % of the activity, suggesting that the disulfide reducing activity of the \(a'\) domain was greater than that of the other domains. The quadruple mutant C36/39/171/174A, in which all four active cysteines were mutated to alanine, had no reducing activity, indicating that cysteines located in the inactive thioredoxin domain, \(b\), were not involved in disulfide reduction.

### 9.3.2 Chaperone Activities of the P5 Mutants

Reduced and denatured lysozyme is commonly used as a substrate to determine whether PDIs have chaperone activity and promote oxidative refolding in vitro. Most PDI family members possess chaperone activity, catalyzing the formation and rearrangement of disulfide bonds during the correct folding of nascent proteins. We have found, however, that P5 possesses anti-chaperone activity, inhibiting the oxidative refolding of the denatured lysozyme (Akama et al. 2010). Western blot analysis has suggested that the anti-chaperone activity of P5 is the result of its intermolecular isomerization with lysozyme, yielding lysozyme aggregates (Akama et al. 2010).

To clarify this mechanism, we compared the ability of P5 and P5 mutants to inhibit oxidative refolding. The \(a\) domain enhanced lysozyme refolding, behaving like thioredoxin alone, whereas the \(a'b\) fragment without the N-terminal \(a\) domain inhibited folding ~30 % compared with the wild type (100 %). The inhibitory activity of equal concentrations of the \(a\) and \(a'b\) fragments was similar to that of \(a'b\) alone, suggesting that the enhancement of refolding by the \(a\) domain did not occur when \(a'b\) was present. The double N-terminal mutant C36/39A had greater inhibitory activity (60 %) than \(a'b\). These results suggested that the presence of the \(a\) domain enhanced the intermolecular isomerization of P5 with denatured lysozyme, and that the covalent bonding of \(a\) to \(a'b\) was essential for complete inhibition. The other N-terminal active domain mutants (C36A, C39A) had ~60 % and the C-terminal active domain mutants (C171A, C174A and C171/174A) had ~75 % of the activity of the wild type, indicating that the \(a\) domain inhibited refolding when combined with the \(a'\) domain.

### 9.3.3 Detection of Lysozyme Aggregates by Western Blotting

Our previous study has indicated that the P5 inhibition of lysozyme refolding is caused by lysozyme aggregates resulting from intermolecular isomerization. Using Western blotting, we found that the quantity of lysozyme aggregates was reduced when refolding was catalyzed by C36A and C171A, compared with native P5 (Fig. 9.2). Spontaneous crosslinking of reduced and denatured lysozyme alone was observed in refolding buffer (Fig. 9.2), explaining why its refolding rate was no higher than 40 %.
In the presence of native P5, lysozyme aggregates were detected in the high molecular region, while lysozyme intermediates were not, similar to our earlier findings (Akama et al. 2010). In the presence of C36A or C171A, lysozyme intermediates had higher molecular weights than those of lysozyme alone, with the amounts of aggregates being smaller than for P5. No aggregates were observed in the presence of a’b, but the sizes of the intermediates were smaller than those observed with the cysteine mutants. These results further indicated that the a and a’ domains cooperatively contributed to the production of lysozyme aggregates and that the a domain accelerated the aggregate production.

9.4 Discussion

9.4.1 Collaborative Isomerization by Two Active Domains

The results of the lysozyme refolding assay showed that N-terminal active cysteines contribute more to the inhibition of refolding than C-terminal active cysteines, similar to findings with rat and human P5 (Kramer et al. 2001; Kikuchi et al. 2002).
Using human P5 variants, a single active domain, particularly the $a$ domain, has been shown sufficient for isomerase activity (Kikuchi et al. 2002). When both cysteines in either the $a$ or $a'$ domain are replaced with serine, the isomerase activities of C36/39S and C171/174S, using insulin as a substrate, are 50% and 75%, respectively, relative to the wild type, suggesting that the $a$ domain can compensate for the replacement of cysteines in the $a'$ domain. However, a study of yeast PDI has shown that the isomerase activity of mutants, in which cysteine residues were replaced with alanine, is less than that of the serine mutants, a finding thought to be the result of the reducing activity remaining in the hydroxyl group of serine (Tian et al. 2006).

Using a lysozyme refolding assay, we found that the inhibitory activity of the N-terminal mutants (C36A, C39A, and C36/39A) was ~60% of that of the wild type, whereas the inhibitory activity of the C-terminal mutants (C171A, C174A, and C171/174A) was ~75%, indicating that both active domains collaborate in intermolecular isomerization. Western blotting showed that comparable amounts of lysozyme aggregates were present in the presence of C36A or C171A, indicating that the $a$ and $a'$ domains cooperatively inhibited refolding. The relatively stronger reducing activity of the $a'$ relative to the $a$ domain observed in the insulin turbidity assay also suggested collaborative isomerization. The refolding inhibitory activity of both $a'b$ and the mixture of $a$ and $a'b$ was 30% of that of the wild type, suggesting that a covalent bond between the two was essential for complete inhibition. This finding was supported by our Western blotting results, with $a'b$ alone showing lysozyme intermediates but no aggregates. The collaborative inhibition by the $a$ and $a'$ domains suggests that they have comparable access to substrates, although these domains are adjacent to each other, possibly because the amino acid chain is relatively long (36 residues) between these domains, enabling the $a$ domain to move widely. The more efficient inhibition by C36/39A, in which both cysteines in the $a$ domain were mutated to inactive alanine, than by $a'b$, which lacked an $a$ domain, suggests that the mobile $a$ domain holds and stabilizes large lysozyme aggregates.

9.4.2 Importance of Thioredoxin Domain Order

Evolution has given rise to the PDI family of proteins, with 1–6 thioredoxin domain(s). The ancestral PDIs may have been characterized by active domains packed together, with these domains structurally separated in descendant PDIs (Pedone et al. 2010). Protein disulphide oxidoreductases (PDOs), composed of only two thioredoxin domains, can be classified as ancestral types. P5 may also be classified as an ancestral PDI because it contains three thioredoxin domains, in the order $a$, $a'$, and $b$, similar to PDOs. The domain structure of P5, consisting of two contiguous active domains and a third inactive domain, may be responsible for its potent isomerization activity and unique function; this in turn may be the result of the nonspecialized roles of both the $a$ and $a'$ domains. Rather, they cooperatively isomerize disulfides. The downregulation of P5 from the epididymal corpus to the cauda suggests that its potent isomerization activity may be involved in sperm maturation.
9.5 Conclusion

We investigated the properties of domain-deleted and cysteine variants of P5 having an active $\alpha$ and $\alpha'$ domain and an inactive $\beta$ domain. The reductive activities of C171A, C174A, and C171/174A ($\alpha'$ domain variants) were ~50% of the wild type, whereas those of C36A, C39A, C36/39A ($\alpha$ domain variants), and $\alpha'\beta$ were ~75%. The inhibitory activities of lysozyme refolding of $\alpha$ and $\alpha'$ domain variants were ~60% and ~75% of wild type, respectively, whereas the inhibitory activity of a mixture of $\alpha$ and $\alpha'\beta$ was 30% as well as that of $\alpha'\beta$. Together with the results of Western blotting, these findings suggested that two active domains of P5 cooperatively inhibited lysozyme refolding by intermolecular isomerization, and that the isomerase activity was enhanced by $\alpha$ domain and the covalent bond between $\alpha$ and $\alpha'\beta$ was essential for the “potent” isomerase activity. Our finding on its isomerization mechanism may contribute for further understanding of molecular basis of sperm maturation, asymmetrical organogenesis, and tumor immune evasion.

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