A Highly Stable D-Amino Acid Oxidase of the Thermophilic Bacterium Rubrobacter xylanophilus

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D-Amino acid oxidase (DAO) is a biotechnologically attractive enzyme that can be used in a variety of applications, but its utility is limited by its relatively poor stability. A search of a bacterial genome database revealed a gene encoding a protein homologous to DAO in the thermophilic bacterium Rubrobacter xylanophilus (RxDAO). The recombinant protein expressed in Escherichia coli was a monomeric protein containing noncovalently bound flavin adenine dinucleotide as a cofactor. This protein exhibited oxidase activity against neutral and basic D-amino acids and was significantly inhibited by a DAO inhibitor, benzoate, but not by any of the tested D-aspartate oxidase (DDO) inhibitors, thus indicating that the protein is DAO. RxDAO exhibited higher activities and affinities toward branched-chain D-amino acids, with the highest specific activity toward D-valine and catalytic efficiency ($k_{cat}/K_m$) toward D-leucine. Substrate inhibition was observed in the case of D-tyrosine. The enzyme had an optimum pH range and temperature of pH 7.5 to 10 and 65°C, respectively, and was stable between pH 5.0 and pH 8.0, with a $T_{50}$ (the temperature at which 50% of the initial enzymatic activity is lost) of 64°C. No loss of enzyme activity was observed after a 1-week incubation period at 30°C. This enzyme was markedly inactivated by phenylmethylsulfonyl fluoride but not by thiol-modifying reagents and diethyl pyrocarbonate, which are known to inhibit certain DAOs. These results demonstrated that RxDAO is a highly stable DAO and suggested that this enzyme may be valuable for practical applications, such as the determination and quantification of branched-chain D-amino acids, and as a scaffold to generate a novel DAO via protein engineering.

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The bacterium *Rubrobacter xylanophilus* DSM9941, first isolated from a thermally polluted runoff from a carpet factory in the United Kingdom in 1996, is a thermophilic actinobacterium with an optimum growth temperature of 60°C (15). This bacterium also exhibits extreme radiation and desiccation resistance. Several enzymes from this thermophilic bacterium, such as mannosyl-3-phosphoglycerate synthase and the enzymes involved in trehalose synthesis (trehalose-6-phosphate synthase, trehalose-6-phosphate phosphatase, and trehalose glycosyltransferase synthase), have been characterized and shown to exhibit higher thermal stability (16, 17), suggesting that this organism may be a candidate source of stable enzymes.

In the present study, we found a gene encoding a DAO homologous protein in the thermophilic bacterium *R. xylanophilus* and show that the gene encodes an active DAO protein with high stability and high affinity for D-amino acids, especially branched-chain D-amino acids. These properties may be valuable for practical applications, such as the determination and quantification of branched-chain D-amino acids, and could make it a good candidate scaffold for developing other useful forms of DAO.

**MATERIALS AND METHODS**

**Materials.** D-Leucine and D-histidine were purchased from Wako Pure Chemical Industries (Osaka, Japan). D-Isoleucine was purchased from the Peptide Institute (Osaka, Japan). D-Aspartate was a generous gift from Tanabe Pharmaceutical (Osaka, Japan). All of the other D-amino acids were purchased from Nacalai Tesque (Kyoto, Japan). Restriction nucleases and other DNA-modifying enzymes were obtained from TaKaRa Bio (Shiga, Japan), Toyobo (Osaka, Japan), or New England Biolabs ( Ipswich, MA). All of the other chemicals were of analytical reagent grade and were purchased from Wako Pure Chemical Industries (Osaka, Japan). D-Aspartate was a generous gift from the Peptide Institute (Osaka, Japan). All of the other chemicals were of analytical reagent grade and were purchased from Wako Pure Chemical Industries, Nacalai Tesque, or Sigma-Aldrich ( St. Louis, MO). Oligonucleotide primers were synthesized by Operon Biotechnology (Tokyo, Japan), and their sequences are listed in Table S1 in the supplemental material.

**Bacterial strains and cultivation media.** *R. xylanophilus* NBR1C100952 (DSM9941) was obtained from the National Institute of Technology and Evaluation (Tokyo, Japan). *R. xylanophilus* cells were grown in NBRC medium no. 802 (1.0% polypeptone, 0.2% yeast extract, 0.1% MgSO₄·7H₂O [pH 7.0]) at 60°C. *Escherichia coli* strain DH5α was used as a host for DNA manipulation, and *E. coli* strain BL21(DE3) or BL21 Star (DE3) was used as a host for the production of RxDAO. *E. coli* cells were cultivated in Luria-Bertani (LB) or Terrific broth (TB) medium (18). When necessary, ampicillin was added to the cultivation media at a concentration of 100 μg/ml.

**Sequencing of the RxDAO gene.** The genomic DNA of *R. xylanophilus* was prepared using a CTAB (cetyltrimethylammonium bromide) procedure (19). Based on the draft genome sequence of *R. xylanophilus* (http://genome.jgi-psf.org/rubxy/rubxy.home.html) and the partial sequence identified in this study, several primers were designed to hydridize to the inside and outside of the putative DAO gene (Rx5_0526) (see Table S1 in the supplemental material). The primer sets RxDAOOp1-2/RDAOOp2N, RxDAOOp1N/RDAOOp2N, RxDASONPCR1/RxDAOOP10, and RxDASONPCR1/RxDAn1R23-43 produced 1.2-, 0.98-, 0.94-, and 0.86-kbp DNA fragments, respectively. PCRs were performed using LA Taq DNA polymerase (TaKaRa Bio), the primer sets were used directly or after cloning into pGEM-T Easy vector or pCR4-TOPO (Invitrogen, Carlsbad, CA). The sequence of the DNA fragment in the vectors was determined using some sequencing primers. Amino acid sequence identities were determined by using the NEEDLE global alignment program EMBoss (http://www.ebi.ac.uk/Tools/psa/emboss_needle/).

**Expression and purification of RxDAO in *E. coli.*** The RxDAO gene was amplified by PCR using LA Taq DNA polymerase, with the RxDAOOp1N and RDA51B15 primers (see Table S1 in the supplemental material) and the bacterial genomic DNA as a template. The amplified fragment was ligated into pGEM-T Easy vector (Promega, Madison, WI), and its sequence was confirmed by sequencing. The plasmid harboring the gene was digested using NdeI and BamHI, and the resulting fragment was ligated into pET15b (Novagen, Madison, WI) to produce the expression vector, pERDAOmHC, in *E. coli.*

*E. coli* BL21(DE3) or BL21 Star(DE3) cells harboring pERDAOmHC were precultivated at 30°C for 14 h in LB medium containing 100 μg of ampicillin/ml, with shaking at 162 rpm. An aliquot of the preculture was transferred to 3 liters of TB medium in a jar fermentor (MDL500-5L; BE Maruhushi, Tokyo, Japan) and cultivated at 37°C. When the optical density at 660 nm reached ~0.5, IPTG (isopropyl-β-D-thiogalactopyranoside) was added to the medium at a final concentration of 0.1 mM, and the cells were further cultivated at 37°C for 13 h. The cells were collected by centrifugation at 5,000 × g for 10 min at 4°C, washed once with anion-exchange (EW) buffer (50 mM sodium phosphate buffer, 0.3 M NaCl [pH 7.0]), and resuspended in the same buffer. The cells were homogenized for 25 s with a sonicator (UD-201; Tomy Seiko, Tokyo, Japan) at an output of 10 and a duty cycle of 50%. The homogenate was centrifuged at 20,000 × g for 30 min at 4°C. The supernatant was incubated at 70°C for 10 min and then centrifuged at 20,000 × g for 30 min at 4°C. After filtration through a 0.45-μm-pore-size membrane filter (Omnipore; Millipore, Billerica, MA), the supernatant was applied to a Talon metal-affinity column (Clontech, Mountain View, CA). The column was washed with EW buffer containing 40 mM imidazole, and bound proteins were eluted using EW buffer containing 150 mM imidazole. The eluted protein solution was concentrated using an Amicon Ultra-15 centrifugal filter device with a 10,000-molecular-weight cutoff membrane (Millipore), and dialyzed overnight against 50 mM potassium phosphate buffer (pH 8.0) containing 5% (vol/vol) glycerol. After centrifugation at 20,000 × g for 30 min at 4°C, the supernatant was stored at ~8°C until use.

**SDS-PAGE and molecular mass determination.** The homogeneity of the purified RxDAO was confirmed by SDS-PAGE using a 12.5% acrylamide gel. The protein bands were visualized by staining with Coomassie brilliant blue R-250. The native molecular mass was determined by gel filtration in a Superdex 200 HR 10/30 column equilibrated with 50 mM potassium phosphate buffer (pH 8.0) containing 150 mM NaCl and 5% (vol/vol) glycerol. The protein sample was loaded and eluted at a flow rate of 0.3 ml/min, with the same buffer. The molecular mass standards used to calibrate the column were aldolase (158 kDa), albumin (67.0 kDa), chymotrypsinogen A (25.0 kDa), and RNase A (13.7 kDa).

**Enzyme assay and protein determination.** Enzyme activity was determined using an oxygen electrode method, a coupled D-α-dianisidine/peroxidase method, or a 2,4-dinitrohydrazine (DNPH) method. In the oxygen electrode method, the reaction mixture contained 20 mM amino acid and 100 μM flavin adenine dinucleotide (FAD) in 50 mM potassium phosphate buffer (pH 8.0). The oxygen consumption in the reaction mixture was monitored using a Clark-type oxygen electrode (Oxygraph system; Hansatech, Norfolk, United Kingdom). In the coupled D-α-dianisidine/peroxidase method, the reaction mixture contained 20 mM amino acid, 0.86 M D-α-dianisidine (Sigma-Aldrich), 25 U of highly stabilized horseradish peroxidase (Sigma-Aldrich), and 20 μM FAD in 50 mM potassium phosphate buffer (pH 8.0) at 60°C. A molar extinction coefficient for oxidized D-α-dianisidine of 8.3 × 10⁴ M⁻¹ cm⁻¹ was used for calculating the enzyme activity. The DNPH assay was performed with D-valine as a substrate in 50 mM potassium phosphate buffer (pH 8.0) at 50°C, as described previously (20). The enzyme activity was determined colorimetrically at 440 nm using a molar extinction coefficient for the ketovaline hydrazone of 1.58 × 10⁴ M⁻¹ cm⁻¹ (21). The protein concentration was determined by using a Bio-Rad protein assay reagent, with bovine serum albumin as the standard.

**Substrate specificity and kinetic parameters.** Substrate specificity was determined for 20 mM each amino acid, except for 2 mM L-tyrosine, in 50 mM potassium phosphate buffer (pH 8.0). The kinetic parameters for D-valine (0.01 to 20 mM), D-leucine (0.01 to 20 mM), D-isoleucine...
RESULTS

Sequence determination of the putative DAO gene of *R. xylanophilus*. To identify DAO homologous proteins in thermophilic prokaryotic organisms, we searched a prokaryotic genome database for the amino acid sequences of ScDAO, RgDAO, and pkDAO. Very limited homology to the DAO sequences was present in the thermophilic and hyperthermophilic bacteria and archaea, but we did identify a DAO homologous protein (Rxyl_0526) with significant homology to the known DAOs in the thermophilic bacterium *R. xylanophilus* DSM9941. To obtain the entire coding gene, we performed PCR using primers that annealed to the 5’- and 3’-terminal sequences derived from the genome sequence. However, despite several attempts under various conditions, we were unable to amplify the gene using this primer set. Several PCR attempts with combinations of various primers annealing within and outside the gene produced some possible DNA fragments of the gene, and the complete sequence of the putative DAO gene was determined by analyzing the PCR fragments.

The nucleotide sequence of the putative DAO gene that we finally determined was somewhat different from that recorded in the genome database: approximately 84 and 82% identities in nucleotide and amino acid sequences, respectively. The gene we isolated had 975 bp coding for a polypeptide of 324 amino acids, with a deduced molecular mass of 35.1 kDa (Fig. 1). A BLAST search revealed that the deduced amino acid sequence of RxDAO had high identities to those of the hypothetical proteins of the Gram-negative bacteria *Stigmatella aurantiaca* DW43-1 (UniProt accession no. Q88XF1, 54%) and *Myxococcus stipitatus* DSM14675 (L7U7T5, 51%) and to those of the Gram-positive bacteria *Streptomyces carleya* DSM46488 (F8JF7T, 47%) and *Kribellia flavida* DSM17836 (D2Q3A8, 49%), all of which are not thermophilic bacteria. The amino acid sequence also exhibited high identities to DAOs: 45.9, 38.0, 31.7, 30.4, 36.1, and 35.8% identities to ScDAO, ApDAO, RgDAO, TvDAO, pkDAO, and human DAO (hDAO) amino acid sequences, respectively. In the amino acid sequence of RxDAO, the residues interacting with the α-carboxy and α-amino groups of the substrate in DAOs were conserved at Y217 and D272, and G299, respectively (24) (Fig. 1 and 6C). In addition, a nucleotide-binding motif (GXGXXG, where “X” is any amino acid) was found in the N-terminal region. However, a peroxisome targeting sequence found in eukaryotic DAOs (25) and the dimerization loop in RgDAO (from S308 to K321) (26) were absent in the C-terminal region, resulting in a lower number of amino acid residues compared to eukaryotic DAOs.

Expression and purification of RxDAO in *E. coli*. To verify whether the gene encodes an active DAO protein, we expressed it in *E. coli* as a histidine-tagged protein. Although most of the recombinant protein expressed at 37°C formed insoluble inclusion bodies, the crude extract of *E. coli* exhibited oxidase activity against d-amino acids. Based on SDS-PAGE, recombinant enzyme in the crude extract was estimated to be ca. 2% of the total protein in the extract. Decreasing the expression temperature to 15°C increased the expression of soluble recombinant protein to ca. 6% but decreased the activity of the crude extract by ca. 10%, although the total expressed protein amount was similar regardless of the expression temperatures of 15 and 37°C.

The recombinant enzyme produced at 37°C was purified (>95% purity) by ~340-fold with a yield of 8.3% from the crude extract of *E. coli* by heat treatment at 70°C for 10 min, followed by...
metal-chelate affinity chromatography (see Table S2 in the supplemental material). The heat treatment increased the total activity by ~1.3-fold. The finally purified RxDAO exhibited a specific activity of 21.1 U/mg for d-valine as a substrate, measured using an oxygen electrode at 60°C. The molecular mass of the purified histidine-tagged enzyme was ~35 kDa, as determined using SDS-PAGE (Fig. 2), whereas the native molecular mass was estimated to be 1 mol per mol of protein, indicating the holo form of the purified enzyme. The dialysis of the enzyme solution against 2 M KBr solution did not release significant amounts of FAD even after 5 days, indicating a noncovalent association of flavin with the protein. TLC analysis of the supernatant showed a single fluorescent spot identical to that of the FAD standard, showing that the prosthetic group of RxDAO is probably FAD (Fig. 3B). The omission of FAD cofactor from the reaction mixture had no significant effect on the enzyme activity, and the cofactor content of the purified enzyme was found to be 1 mol per mol of protein, indicating the holo form of the purified enzyme. The dialysis of the enzyme solution against 2 M KBr solution did not release significant amounts of FAD even after 5 days, indicating a tight cofactor binding.

Cofactor analysis. To determine whether the purified protein contains flavin, we analyzed its absorption spectrum (Fig. 3A). The purified protein solution was yellow, and it exhibited a typical absorption spectrum of flavoproteins, with maxima at 273, 370, and 455 nm with an $A_{273}/A_{455}$ ratio of 8.61. The addition of $d$-valine into the protein solution bleached the absorption maxima, indicating a reduction in flavin content. The supernatant of the thermally denatured protein was also yellow with a flavin absorption spectrum, indicating a noncovalent association of flavin with the protein. TLC analysis of the supernatant showed a single fluorescent spot identical to that of the FAD standard, showing that the prosthetic group of RxDAO is probably FAD (Fig. 3B). The omission of FAD cofactor from the reaction mixture had no significant effect on the enzyme activity, and the cofactor content of the purified enzyme was found to be 1 mol per mol of protein, indicating the holo form of the purified enzyme. The dialysis of the enzyme solution against 2 M KBr solution did not release significant amounts of FAD even after 5 days, indicating a tight cofactor binding.

FIG 1 Comparison of amino acid sequences of RxDAO with those of DAOs from other organisms. The amino acid sequences of Rhodobacter xylophilus (Swiss-Prot accession no. P80324), pig kidney (Swiss-Prot accession no. P00371), human (Swiss-Prot accession no. P14920), Trigonopsis variabilis (UniProt accession no. Q99042), Arthrobacter protophormiae (UniProt accession no. Q7X2D3), and Streptomyces coelicolor (UniProt accession no. Q9X7P6) DAOs were aligned using CLUSTAL W. A nucleotide-binding motif (GXGXXG, where “X” is any amino acid) is indicated by closed circles. The functionally important amino acid residues of DAO are indicated by closed triangles. The dimerization loop in RgDAO is indicated by a wavy line. The active-site residues involved in the oxidative inactivation of RgDAO and TvdDAO are indicated by open triangles. The positions of cysteine residues of RxDAO are indicated by open circles. The positions of amino acid residues H-bonded to the flavin O-2 atom are indicated by open squares. The positions of amino acid residues H-bonded to the flavin O-3 atom are indicated by open diamonds.
Substrate specificity and kinetic parameters. We analyzed the relative activities of purified RxDAO toward various amino acids (Table 1). The enzyme exhibited higher oxidase activity toward neutral D-amino acids but lower or no activity toward basic and acidic D-amino acids and L-amino acids. The highest activity was observed for D-valine, with decreasing order of activity for D-leucine, D-isoleucine, and D-tyrosine. These results clearly show that the enzyme is DAO.

The enzyme displayed typical Michaelis-Menten kinetics for the substrates, except for D-tyrosine, which caused substrate inhibition (see Fig. S1 in the supplemental material). The highest catalytic rate constant $k_{\text{cat}}$ was presumed to be for D-tyrosine, followed by D-valine, D-leucine, and D-isoleucine, in decreasing order (Table 2). On the other hand, the lowest Michaelis constant $K_m$ was obtained for D-leucine, followed by D-isoleucine, D-valine, and D-tyrosine, in increasing order. As a result, the enzyme showed the highest catalytic efficiency ($k_{\text{cat}}/K_m$) toward D-leucine, followed by D-isoleucine and D-valine.

Effect of pH and temperature on the activity and stability. The enzyme showed higher activity in the alkaline pH range of 7.5 to 10 and was stable at a pH of 5.0 to 8.0 for 1 h at 55°C (Fig. 4A). The enzyme displayed maximum activity at 65°C (Fig. 4B), which is in good agreement with the optimum growth temperature of R. M1.
*xylanophilus* and similar to that of other enzymes from this bacterium (15, 16). An Arrhenius plot showed that the activation energy of the reaction was approximately constant in the temperature range from 30°C to 50°C, and the activation energy was calculated as 62.1 kJ/mol (see Fig. S2 in the supplemental material). It was stable up to 60°C at a protein concentration of 1.0 mg/ml, with a *T* 50 value (the temperature at which 50% of initial enzymatic activity is lost) of ca. 64°C (Fig. 4B). This thermal stability was, however, dependent on the enzyme protein concentration, and the addition of FAD markedly protected the enzyme from the inactivation (Fig. 4C).

The long-term stability of the enzyme was assessed at 30 and 50°C. Incubation at 30°C caused no significant loss of activity after 1 week, and ca. 40% of the initial activity was observed even after 28 days (Fig. 4D). Incubation at 50°C linearly decreased the activity from that observed in the beginning, but 25% of the initial activity remained after 8 days of incubation. The half-lives of the enzyme were calculated to be 462 h at 30°C and 114 h at 50°C.

**Effect of inhibitors on the enzyme activity.** The inhibitory effect of known DAO (benzoate, anthranilate, and crotonate) and DDO (malonate, *meso*-tartrate, and *d*-malate) inhibitors, and amino acid-modifying reagents on RxDAO was examined (Table 3). Among DAO inhibitors, benzoate strongly inhibited the enzyme activity (15, 16). An Arrhenius plot showed that the activation energy of the reaction was approximately constant in the temperature range from 30°C to 50°C, and the activation energy was calculated as 62.1 kJ/mol (see Fig. S2 in the supplemental material). It was stable up to 60°C at a protein concentration of 1.0 mg/ml, with a *T* 50 value (the temperature at which 50% of initial enzymatic activity is lost) of ca. 64°C (Fig. 4B). This thermal stability was, however, dependent on the enzyme protein concentration, and the addition of FAD markedly protected the enzyme from the inactivation (Fig. 4C).

![Graph A](https://via.placeholder.com/150)

**FIG 4** Effect of pH, temperature, protein concentration, and FAD on the enzyme activity and stability of RxDAO. (A) Effect of pH on the enzyme activity (○) was determined by measuring the activity at the indicated pH values at 55°C. Effect of pH on the enzyme stability (●) was determined by measuring the remaining activity at pH 8.0 after incubation of the enzyme (0.1 mg/ml) at various pH values at 50°C for 1 h. (B) Effect of temperature on the enzyme activity (○) was determined by measuring the activity at the indicated temperatures at pH 8.0. Effect of temperature on the enzyme stability (●) was determined by measuring the remaining activity after incubation of the enzyme (0.1 mg/ml) at the indicated temperatures for 1 h. (C) Effect of enzyme protein concentration on the enzyme activity was determined by measuring the remaining activity after incubation at 60°C for 1 h. The gray bar represents the effect of 100 μM FAD on thermal denaturation. (D) The long-term thermal stability of the enzyme was determined by measuring the activity during the incubation of the enzyme (0.1 mg/ml) at 30°C (●) and 50°C (▲) at pH 8.0. The assays for pH, thermal, and long-term thermal stabilities and the effect of the enzyme concentration were performed using a coupled *o*-dianisidine/peroxidase method at 60°C, and optimum pH was determined using a DNPH method at 55°C. The assay for optimum temperature was performed using an oxygen electrode. Each data point represents the mean ± the standard deviation of three measurements.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Mean relative activity (% ± SD)</th>
</tr>
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<tbody>
<tr>
<td>None</td>
<td>100</td>
</tr>
<tr>
<td><strong>DAO inhibitors</strong></td>
<td></td>
</tr>
<tr>
<td>Benzoate</td>
<td>37.0 ± 13.0</td>
</tr>
<tr>
<td>Anthranilate</td>
<td>83.3 ± 5.6</td>
</tr>
<tr>
<td>Crotonate</td>
<td>89.2 ± 3.3</td>
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<tr>
<td><strong>DDO inhibitors</strong></td>
<td></td>
</tr>
<tr>
<td>Malonate</td>
<td>99.0 ± 2.7</td>
</tr>
<tr>
<td><em>meso</em>-Tartrate</td>
<td>99.5 ± 16.0</td>
</tr>
<tr>
<td><em>d</em>-Malate</td>
<td>107 ± 16</td>
</tr>
<tr>
<td><strong>Amino acid-modifying reagents</strong></td>
<td></td>
</tr>
<tr>
<td>DTNB</td>
<td>87.4 ± 5.5</td>
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<tr>
<td>Iodoacetic acid</td>
<td>112 ± 6</td>
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<tr>
<td>PMSF</td>
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</tr>
<tr>
<td>2,3-Butanediol</td>
<td>115 ± 3</td>
</tr>
<tr>
<td>DEPC</td>
<td>90.2 ± 1.1</td>
</tr>
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</table>

*For DAO and DDO inhibitors, the enzyme activity was assayed in the presence of a 10 mM concentration of each compound. For amino acid-modifying reagents, the enzyme (5 μg/ml) was incubated with a 5 mM concentration of each compound at 30°C for 30 min, and the remaining activity was measured. The assays were performed by the DNPH method with 20 mM *o*-valine as a substrate at 55°C. Each value represents the mean of triplicate measurements.*
zyme activity by 63%, whereas the others weakly inhibited the enzyme activity. In contrast, none of the DDO inhibitors inhibited the enzyme. Among amino acid-modifying reagents, the serine residue-modifying reagent phenylmethylsulfonyl fluoride (PMSF) strongly inhibited the enzyme activity by 68%, and the cysteine- and histidine-modifying reagents 5,5'-dithiobis (2-nitrobenzoic acid) and diethyl pyrocarbonate (DEPC) weakly inhibited the enzyme activity. In contrast, the cysteine- and arginine-modifying reagents, iodoacetic acid and 2,3-butanedione, did not inhibit the enzyme activity. These results suggest that a serine residue may be particularly important for the catalytic activity of the enzyme.

**DISCUSSION**

In this study, we showed that a DAO homologous gene in the thermophilic bacterium *R. xylanophilus* encodes an active DAO protein that has high stability and high activity and affinity for branched-chain D-amino acids. These properties may be valuable for practical applications, particularly for the detection and quantification of branched-chain D-amino acids. RxDAO could also be a promising scaffold for developing novel DAOs that are useful for other practical purposes.

The nucleotide sequence of the RxDAO gene is somewhat different from the sequence from the genome database. Such differences have also been reported for other genes from *R. xylanophilus* (16), indicating the necessity for reanalyzing the genome sequence. Interestingly, the amino acid sequence of RxDAO showed a higher identity to hypothetical proteins of the phylogenetically distant Gram-negative bacteria *S. aurantia* and *M. stipitatus* than that of DAOs of phylogenetically close Gram-positive bacteria *S. coelicolor* and *A. protophormiae*. Phylogenetic analysis also confirms that RxDAO forms a distinct subgroup with the putative DAO proteins of Gram-negative bacteria, but not with DAOs and putative DAOs of Gram-positive bacteria, including *R. radiotolerans* that belongs to the same genera, suggesting a close phylogenetic relationship between them (Fig. 5).

The fermentation yield of RxDAO in *E. coli* was ∼4.0 U/g cell (see Table S2 in the supplemental material). This yield was comparable to that of human DAO under an unoptimized condition but 10- to 230-fold lower than those of microbial DAOs (6, 27). Furthermore, the amount of soluble RxDAO protein in the crude extract of recombinant cells was similar to that of ScDAO but lower than those of other microbial DAOs (6). The increase in the yield by heat treatment suggests that a part of the expressed recombinant protein might be trapped as a soluble folding intermediate (see Table S2 in the supplemental material), which may be due to the lower expression temperature than that in the original host. Similar to some other DAOs, RxDAO was produced as an insoluble protein in *E. coli*. However, as shown in the production of recombinant hDAO and RgDAO, further optimization of the expression conditions could improve the production of the active enzyme (27, 28).

Although most DAOs are dimers, RxDAO is a monomer. DAOs of rat (rDAO) and the yeast *Candida boidinii* (CbDAO) are also monomers, but the former enzyme forms a dimer in the presence of an inhibitor (29, 30). In addition, the oligomerization state of pkDAO is dependent on protein concentration (2). However, the oligomerization state of RxDAO in the presence of inhibitor and its dependence on the enzyme protein concentration are not known. Two types of dimer formation of DAO have been reported, namely, head-to-tail and head-to-head dimers (2). RgDAO forms a head-to-tail dimer, in which a long C-terminal loop, which is absent from RxDAO, as well as other DAOs, is responsible for the dimer formation (Fig. 1). Mammalian DAOs form a head-to-head dimer, and the affinity of this interaction is considered to be related to the electrostatic surface potential at the dimer interface: a more positively charged surface is likely to interact more weakly (31). In a three-dimensional model of RxDAO (Fig. 6), analogous to that of pkDAO and RgDAO, the corresponding surface appears to be positively charged (Fig. 6B). These results suggest a possible mechanism for the monomer formation of RxDAO.

The substrate specificity of RxDAO is similar to that of ScDAO, which prefers branched-chain D-amino acids, but largely different from that of the other bacterial oxidase ApDAO (4, 5). RxDAO showed only a very low activity toward D-alanine and D-methionine, which are preferred substrates for most eukaryotic DAOs and yeast DAOs, respectively (10). It has recently been shown that branched-chain D-amino acids are produced by various bacterial species and are suggested to be involved in cell wall remodeling and biofilm dispersion (32). Although the production of branched-chain D-amino acids remains unknown in *R. xylanophilus*, the substrate specificity indicates the possibility that RxDAO might be involved in the functions of branched-chain D-amino acids through their degradation. In pkDAO, a flexible loop near the active site, called the active-site lid that opens and closes for substrate entry and product exit, is involved in substrate specificity (33, 34). In particular, the flexibility of Y224 in the lid is likely important for its broad substrate specificity. In the active-site model of RxDAO, L215 within the active-site lid region D208-Y217 appears to be located in the corresponding spatial position of Y224 (Fig. 1 and 6C). This difference might contribute to the different substrate specificity. In addition to L215, the ligand side chain is likely to be covered with hydrophobic amino acid residues, namely, V49, Y51, Y53, and L207 (Fig. 6C). These hydrophobic amino acid residues possibly also contribute to the substrate specificity of RxDAO.

The specific activity of RxDAO for D-valine was much lower than that of the microbial DAOs, ApDAO (180 U/mg for D-methionine), RgDAO (175 U/mg for D-alanine), TvDAO (86.6 U/mg for D-alanine), and CbDAO (139 U/mg for D-alanine) but was similar to or higher than that of the mammalian DAOs, hDAO (18 U/mg for D-proline) and pkDAO (8.8 U/mg for D-alanine) (http://www.brenda-enzymes.info). It should be noted, however, that a decreased dissolved oxygen concentration in the reaction mixture might affect the activity of RxDAO. In contrast to the specific activity, the binding affinity of RxDAO for branched-chain D-amino acids was from 1 to 2 orders of magnitude higher than that of other DAOs, including the bacterial ApDAO (Table 2). This property might be valuable, in particular for the determination and quantification of branched-chain D-amino acids. RxDAO could contribute to clarifying the physiological function of the D-amino acids in bacteria by providing a means to quantify branched-chain D-amino acids. RxDAO was, however, subject to substrate inhibition by D-tyrosine (Table 2; see also Fig. S1 in the supplemental material). Such inhibition was also observed in hDAO by D-tyrosine, D-phenylalanine, and D-DOPA (35). The inhibitory mechanism has not yet been determined but is suggested to be
Gram-negative bacteria

Animal DAOs

Fungal DAOs

![Phylogenetic tree of DAOs and bacterial homologous proteins](image)

**FIG 5** Phylogenetic tree of DAOs and bacterial homologous proteins. The phylogenetic analysis was performed using the neighbor-joining method in MEGA6.0 with 1,000 bootstrap trials. The numbers at nodes indicate bootstrap value percentages. The asterisks indicate putative bacterial DAOs. The accession numbers of the amino acid sequences used for the analysis were as follows: R. gracilis (Swiss-Prot accession no. F80324), pig (Swiss-Prot accession no. P00371), human (Swiss-Prot accession no. P14920), mouse (Swiss-Prot accession no. P18894), Pichia pastoris (UniProt accession no. C4R4G9), Candida boidinii (UniProt accession no. Q9HG3), Schizosaccharomyces pombe (UniProt accession no. Q9Y7N4), T. variabilis (UniProt accession no. Q99042), carp (UniProt accession no. A8WAD5), nematode (UniProt accession no. Q95XG9), Micromonospora aurantiaca (UniProt accession no. D9T33), Actinoplanes friulensis (UniProt accession no. U5W5C2), Kribbella flavida (UniProt accession no. D2Q3A8), Myxococcus xanthus (UniProt accession no. L7U7T5), Stigmatella aurantiaca (UniProt accession no. Q8XSF1), Actinosynnema missouriensis (UniProt accession no. G6WHB6), Geodermatophilus obscurus (UniProt accession no. D5U2R), A. protophormiae (UniProt accession no. Q7X2D3), Tsukamurella paurometabola (UniProt accession no. D5U2R), Isoperictola variabilis (UniProt accession no. F6FRQ3), Nocardioideas sp. (UniProt accession no. A1SHK8), S. coelicolor (UniProt accession no. Q9Y7P6), Streptomyces scabies (UniProt accession no. CSZB51), Streptomyces pratenis (UniProt accession no. EW8AE4), Streptomyces albicus (UniProt accession no. D6B010), Streptomyces cattleya (UniProt accession no. F8JEF7), Actinoplanes missouriensis (UniProt accession no. 1O02Z4), Stackebrandtia nassauensis (UniProt accession no. D3Q6B1), Mycobacterium avium (UniProt accession no. A0QGE5), Mycobacterium kansasii (UniProt accession no. U5W1K4), Mycobacterium tuberculosis (UniProt accession no. P9WP27), Nocardia brasiliensis (UniProt accession no. K0F255), Saccharopolyspora erythraea (UniProt accession no. A4F8D6), Rubrobacter radiotolerans (NCBI-GI accession no. 62778579), Streptomyces albulus (NCBI-GI accession no. 636570506), and Kutzneria albia (NCBI-GI accession no. 578013459).

such that a second substrate binds to the active site in an orientation that makes the progression of the reaction unlikely.

The importance of a serine residue in the catalytic activity of RxDAO is suggested by the inhibition by PMSF (Table 3). RxDAO has a serine residue (S300) that is located next to the glycine residue that possibly interacts with the substrate (Fig. 1 and 6C), suggesting its possible involvement in the inactivation. In contrast, thiol-modifying reagents, which can strongly inhibit many DAOs, did not significantly inhibit RxDAO. Such reagents inactivate TvDAO and RgDAO by the oxidation of C298 and C208, respectively (36, 37). However, no cysteine residue is conserved at the corresponding positions of RxDAO or is found near the active site. In addition, DEPC, which can markedly inhibit TvDAO and pkDAO, did not significantly inhibit the enzyme activity. pkDAO is inactivated by the modification of H217 in the active-site lid (38). RxDAO has no histidine residue in the lid or in the active site (Fig. 1).

The pH-stability profile of RxDAO appeared similar to the profiles of RgDAO, TvDAO, and ApDAO but not to the profiles of pkDAO and ChDAO, which are stable under alkaline conditions (6, 9). The pH-activity profile of RxDAO was similar to the profiles of RgDAO and pkDAO in terms of higher activity at alkaline pHs but not to the profiles of TvDAO, ChDAO, and ApDAO, which show lower activity at pH values above 8.5 (39). RxDAO exhibited a much higher optimum temperature and $T_{50}$ value than TvDAO that had the highest optimum temperature (55°C).
and thermal stability (approximately 52°C of $T_{50}$ value) among known DAOs thus far (14), indicating a much higher stability of the enzyme. The protein concentration-dependent stability observed for RxDAO has also been reported for TvDAO and RgDAO but not for pkDAO (6,10). The protective effect of FAD against enzyme inactivation observed for RxDAO has also been reported in RgDAO, in which the release of FAD causes protein denaturation and a looser conformation in the yeast enzyme (40,41). Therefore, one reason for the higher stability of RxDAO might be its higher affinity for the cofactor, as suggested based on the resistance of flavin to dissociate from the complex. The environment of the flavin O-2 position is one of the factors involved in the affinity of DAO for the cofactor (42). In pkDAO, the flavin O-2 atom is H bonded to T317, whereas in RgDAO, the atom is tightly bound to Y338 and Q339, resulting in the higher flavin-binding affinity of RgDAO (43). Similar to pkDAO, RxDAO is likely to have V302 and T303 in the corresponding positions, respectively (Fig. 1), suggesting that other factors are likely responsible for the higher affinity. Further studies, such as the determination of the crystal structure, may be required to reveal the mechanism underlying the higher stability of RxDAO.

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REFERENCES

