

Serine racemases from barley, *Hordeum vulgare* L., and other plant species represent a distinct eukaryotic group: Gene cloning and recombinant protein characterization [☆]

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Abstract

Several D-amino acids have been identified in plants. However, the biosynthetic pathway to them is unclear. In this study, we cloned and sequenced a cDNA encoding a serine racemase from barley which contained an open reading frame encoding 337 amino acid residues. The deduced amino acid sequence showed significant identity to plant and mammalian serine racemases and contained conserved pyridoxal 5-phosphate (PLP)-binding lysine and PLP-interacting amino acid residues. The purified gene product catalyzed not only racemization of serine but also dehydration of serine to pyruvate. The enzyme requires PLP and divalent cations such as Ca^{2+} , Mg^{2+} , or Mn^{2+} , but not ATP, whereas mammalian serine racemase activity is increased by ATP. In addition to the results regarding the effect of ATP on enzyme activity and the phylogenetic analysis of eukaryotic serine racemases, the antiserum against *Arabidopsis* serine racemase did not form a precipitate with barley and rice serine racemases. This suggests that plant serine racemases represent a distinct group in the eukaryotic serine racemase family and can be clustered into monocot and dicot types.

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1. Introduction

Serine racemase, which catalyzes racemization of L- and D-amino acids, is found in mammals as a biosynthetic enzyme for D-serine (**1a**), with the latter acting as an agonist at the glycine site of the N-methyl-D-aspartate receptor in the mammalian nervous system (Dunlop et al., 1986; Nagata et al., 1989; Hashimoto et al., 1992, 1993, 1995; Nagata et al., 1994). Mammalian serine racemase is a unique enzyme that catalyzes not only serine (**1a**, **1b**) race-

mization but also dehydration of **1a** and **1b** to pyruvate (**2**) in the presence of divalent cations; this distinguishes it from bacterial amino acid racemases (Cook et al., 2002; De Miranda et al., 2002; Strisovsky et al., 2003; Foltyn et al., 2005) (Fig. 1). In plants, several D-amino acids have been discovered in pea seedlings (Ogawa et al., 1977), barley grain, hops blossoms (Erbe and Bruckner, 2000), and tobacco leaves (Kullman et al., 1999). Recently, a serine racemase gene was cloned from *Arabidopsis thaliana* and expressed to produce a protein that has both racemase and dehydratase activities (Fujitani et al., 2006). This gene also has 46% and 45% identity to human and mouse serine racemases, respectively, and the pyridoxal 5-phosphate (PLP)-binding domain is conserved as with mammalian serine racemases. The PLP-dependent *A. thaliana* serine racemase also requires divalent cations but not ATP, whereas there is synergy between Mg^{2+} and ATP in the

Abbreviations: PLP, pyridoxal 5-phosphate; PCR, polymerase chain reaction; IPTG, isopropyl-β-D-thiogalactopyranoside.

[☆] The sequence reported in this paper has been deposited in the DDBJ, EMBL, and GenBank data banks under the accession no. [AB271213](https://www.ncbi.nlm.nih.gov/nuclot/AB271213).

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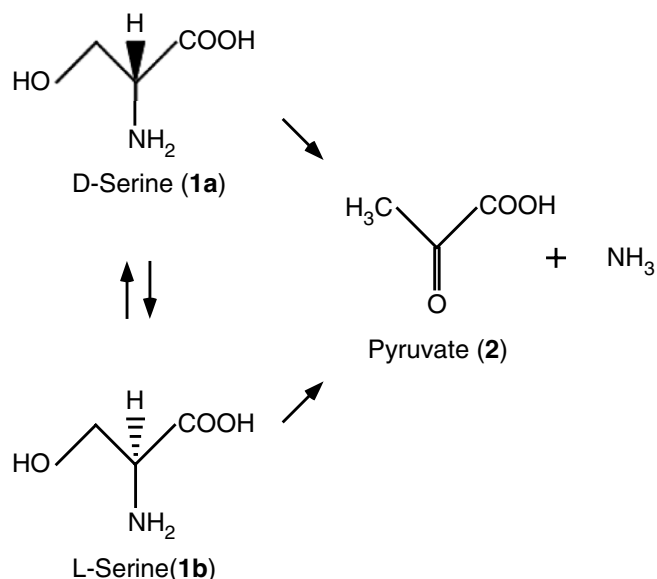


Fig. 1. Reaction pathways of D-serine (**1a**) and L-serine (**1b**) by mammalian serine racemase.

activity of mammalian serine racemases. These reports suggest that this plant serine racemase is distinct from mammalian serine racemases. However, there has been no report on the cloning and characterization of plant serine racemases except for that in *A. thaliana*.

In this report, we describe the isolation and characterization of a cDNA encoding serine racemase from barley. In addition, we obtained a rice serine racemase by cloning a cDNA encoding a homolog of barley serine racemase and compared the structure–function relationships of plant and mammalian serine racemases.

2. Results

2.1. Sequence analysis of barley serine racemase

The resulting nucleotide sequence of the barley serine racemase cDNA, except for the poly(A) sequence, is 1380 bp in length and contains an open reading frame encoding a polypeptide of 337 amino acid residues with a calculated molecular mass of 35.7 kDa (HvSR, AB271213). The deduced amino acid sequence of HvSR showed 89% identity with that of a putative rice serine racemase in data banks (OsSR, AL606647), and 68%, 46%, 45%, and 28% identity with a serine racemase from *A. thaliana* (AtSR, AB206823, Fujitani et al., 2006), as well as human (hSR, AF169974, De Miranda et al., 2000) and mouse (mSR, AF148321, Wolosker et al., 1999) serine racemases, and the catalytic domain of L-threonine dehydratase from *Escherichia coli* (EcTD, Swiss-Prot no. P04969, Gallagher et al., 1998), respectively. The amino acid sequence alignment (Fig. 2) shows that plant and mammalian serine racemases belong to the fold type II PLP enzymes. Specifically, it is known that Lys62 binds

to PLP, Phe61 and Gly241 sandwich the PLP ring, the side-chain of Ser315 is hydrogen-bonded to the pyridium nitrogen of PLP, and Asn89 stabilizes the 3' oxygen of PLP by hydrogen bonding. Additionally, in the PLP-binding domain of *E. coli* threonine dehydratase, a typical fold type II PLP enzyme, these residues correspond to Lys67, Phe66, Gly246, Ser322, and Asn94 of HvSR, respectively, moreover, all of these residues were conserved in rice, *A. thaliana*, human, and mouse serine racemases. The glycine-rich loop, which coordinates the phosphate group of PLP, is composed of a triglycine loop (Gly194–196 of HvSR) in the three plant serine racemases known, whereas it is a tetraglycine loop in human and mouse serine racemases. The phylogenetic tree presented shows that plant serine racemases represent a distinct group from mammalian serine racemases, and that the monocot serine racemases of barley and rice are distinct from the dicot serine racemase of *A. thaliana* (Fig. 3).

2.2. Purification and enzyme activity of recombinant barley serine racemase

SDS-PAGE analysis of *E. coli* cells harboring pHvSR contained an extra protein with the expected molecular mass of ca. 36 kDa, the expression level of which reached ca. 10% of the total protein after 8 h of isopropyl- β -D-thiogalactopyranoside (IPTG) induction. Purification of recombinant HvSR by Ni-NTA column chromatography yielded an ~95% pure preparation (Fig. 4). The purified HvSR was incubated with 5 μ M PLP and 1 mM CaCl_2 which activates AtSR and mammalian serine racemases (Cook et al., 2002; De Miranda et al., 2002; Strisovsky et al., 2003), and the formation of **1a** and **2** was determined. The specific activities for racemization of L-serine (**1b**) to **1a**, and dehydration of **1b** to **2**, were 14 and 228 nmol/min mg, respectively, and those for racemization of **1a** to **1b**, and dehydration of **1a** to **2**, were 2.2 and 89 nmol/min/mg, respectively.

The purified rice serine racemase expressed in *E. coli* cells harboring pOsSR had a molecular mass ~36 kDa, as estimated by both the deduced amino acid sequence and SDS-PAGE, respectively. It was also purified by Ni-NTA column chromatography (Fig. 4), and displayed racemase and dehydratase activities for **1b** of 10 and 214 nmol/mg min, respectively, and those for **1a** of 1.5 and 77 nmol/mg min, respectively.

2.3. Enzymatic characterization of recombinant barley serine racemase

A maximum at 420 nm in the absorption spectrum of the purified HvSR disappeared after dialysis against a buffer containing 10 mM sodium borohydride with concomitant loss of enzyme activity (Fig. 5). Additionally, the enzyme activity of HvSR was inhibited completely by 0.1 mM hydroxylamine (data not shown), indicating that

HvSR	—MGRDDDGHSSTEGGG—AADINSIREARARIAPYVHKTPILSSSTSIDAIAQKQFFKCECFQKAGAFKIRGA	71
OsSR	MGRSGSGDGAESHG—AADIHSIREAQARIAPYVHKTPVLSSTSIDAIVGKQFFKCECFQKAGAFKIRGA	72
AtSR	-----MEANREKYAADILSIKEAHDRIKPYIHRTPVLTSESLNSISGRSLFFKCECFQKAGAFKIRGA	63
hSR	-----MCAQYICISFADVEKAHINIRDSITLTPVLTSSILNQLTGRNFFKCECFQKAGAFKIRGA	60
mSR	-----MCAQYICISFADVEKAHINIQDSITLTPVLTSSILNQLTGRNFFKCECFQKAGAFKIRGA	60
EcTD	-----MADSQPLSGAPEGAEYLRAVLRAVYEAAGVTPLQKMEKLSRLDNVILVKREDROPVHSEFKIRGA	66
HvSR	SNSIFALD—DSQAAKGVVTHSSGNHAAVALAALKRGIPAYIVIPKNAPACKVENVRRYGGQVWSDVTM	140
OsSR	SNSIFALD—DDEASKGVVTHSSGNHAAVALAALKRGIPAYIVIPKNAPACKVDNVKRYGGHIWSDVSI	141
AtSR	GNVLSLD—AEQAAKGVVTHSSGNHAAVALAALKRGIPAYIVIPKNAPACKVDNVIRYGGKVIWSEATM	132
hSR	LNAVRSLVPDALERKPKAVVTHSSGNHGAALTYAAKLEGIPAYIVVPQAPDCKKLAIQAYGASIVYCEPSD	132
mSR	LNAIIRGLIPDTPEEKPKAVVTHSSGNHGAALTYAAKLEGIPAYIVVPQAPDCKKLAIQAYGASIVYCDPSD	132
EcTD	YAMMAGLLE—EQKAHGVITIASAGNHAQGVAFSSARLGVKALIVMPTATADIVKDAVRGFGGEVLLHGANF	135
HvSR	ESRESIAKKVQEEETGAILIHPFNDKTYISGGGTIVCLELLEQVPEIDTIIIVPISGGGLISGVTLAAKAINPSI	212
OsSR	ESRESIAKKVQEEETGAILIHPFNDKTYISGGGTIVCLELLEQVPEIDTIIIVPISGGGLISGVTLAAKAINPSI	213
AtSR	SSREEIASKVLQETGSLIHPYNDGRIISGGGTIALLELEQIQTDAIVVPISGGGLISGVTLAAKAINPSI	204
hSR	ESRENIAKKVTEETEGIMVHPNQEPAVIAGGGTIALLELVNQVPLVDALVVPVGGGMLAGIATVKALKPSV	204
mSR	ESREKVTQRIMQETEGILVHPNQEPAVIAGGGTIALLELVNQVPLVDALVVPVGGGMLAGIATVKALKPSV	204
EcTD	DEAKAKAIELSQQGGFTWVPPFDHPMVVAGGGTIALLELVNQVPLVDALVVPVGGGMLAGIATVKALKPSV	207
HvSR	RILAAEPKGADDSAQSKAAGRIIK—LPATSTIADGLRAF—LGDLTWPVVRDLVDDVIVVDDNATVDAMKMCY	236
OsSR	RILAAEPKGADDSAQSKAAGRIIK—LPSTNTIADGLRAF—LGDLTWPVVRDLVDDVIVVDDNATVDAMKMCY	283
AtSR	RILAAEPKGADDSAQSKAAGRIIK—LPVTNTIADGLRAS—LGDLTWPVVRDLVDDVIVVDDNATVDAMKMCY	274
hSR	KVYAAEPSNADDCYQSKLKGKLMPLNLYPETIADGVKSS—IGLNTWPIIRDLVDDVFTVTEDEIKCATQLVW	275
mSR	KVYAAEPSNADDCYQSKLKGKLMPLNLYPETIADGVKSS—IGLNTWPIIRDLVDDVFTVTEDEIKCATQLVW	275
EcTD	KVIAVLEAEDS—ACLKAALDAGHPVD—PRVGLFAEGVAVKRIQDEIFRLCQEYLDIIITVDSDAICAMKDLF	278
HvSR	ETLKVAEPPSGAIGLAALSDDEFKQSSAWHESSKIGIIVSGGNVDLRVLWDSLYK-----	337
OsSR	EMLKVAEPPSGAIGLAALSDDEFKQSSAWHESSKIGIIVSGGNVDLRVLWDSLYK-----	339
AtSR	EILKVSVEPPSGAIGLAALSDDEFKQSSAWHESSKIGIIVSGGNVDLRVLWDSLYK-----	331
hSR	ERMKLLIEPTAGVGAVALSQHQTQVS—PEVKNIQIVLSSGNVDLTSSITWVKQAERPASYQSVSV	340
mSR	GRMKLLIEPTAGVGAVALSQHQTQVS—PEVKNIQIVLSSGNVDLTSSITWVKQAERPASYQSVSV	339
EcTD	EDVRAVAEPPSGALALAG—MKKYIALHN—IRGERLAHILSGANV-----	320

Fig. 2. Alignment of the deduced amino acid sequences of barley and rice serine racemases with those of *A. thaliana*, human, and mouse serine racemases, as well as *E. coli* biosynthetic L-threonine dehydratase. Barley, rice, *A. thaliana*, human, and mouse serine racemases (HvSR, OsSR, AtSR, hSR, and mSR) correspond to GenBank accession nos. AB271213, AL606647, AB206823, AF169974, and AF148321, respectively, whereas the *E. coli* biosynthetic L-threonine dehydratase (EcTD) corresponds to Swiss-Plot accession no. P04968. Gaps, indicated by dashes, are introduced in the sequences to maximize the homology. Identical amino acid residues among either serine racemases or all six of the enzymes are represented by black boxes. The closed arrowhead, open arrowheads, and asterisks indicate the PLP-binding Lys, amino acid residues that interact with PLP, and the glycine-rich loop, respectively.

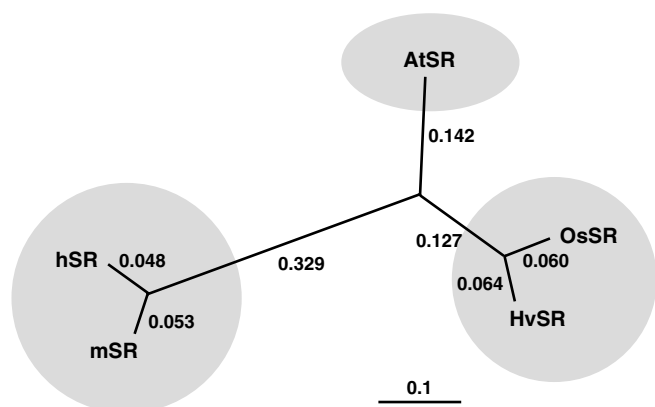


Fig. 3. Phylogenetic analysis of eukaryotic serine racemases. The scale bar represents 0.1 amino acid substitutions per site.

HvSR depends on PLP and is bound to PLP through a Schiff base (Grishin et al., 1995).

The optimum pH and concentration of CaCl_2 in the racemization and dehydration of **1b** were examined in a reaction mixture containing 10 μM PLP. HvSR has maximum activities for both racemization and the elimination reaction at pH 8.5 in the presence of 1 mM CaCl_2 .

The racemase activities for converting various **1b** to **1a** acids were examined in a reaction mixture containing 1 mM CaCl_2 at pH 8.5. HvSR showed high substrate specificity with regard to **1b**, 14 nmol/mg min, whereas L-alanine, L-glutamine, and L-arginine were poorer substrates: the activities were 1.9, 0.74, and 1.7 nmol/mg min, respectively. The K_m and V_{\max} were estimated in a reaction mixture containing 1 mM CaCl_2 , 10 μM PLP at pH 8.5, with corresponding values for converting **1b** to **1a** being 2.6 mM and 21 nmol/mg min, and for **1a** to **1b** of 9.0 mM and 455 nmol/mg min, respectively. Additionally, the K_m and V_{\max} for dehydratase activity from **1b** to **2** were 2.7 mM and 2.8 nmol/mg min, whereas those of **1a** to pyruvate were 8.3 mM and 161 nmol/mg min, respectively.

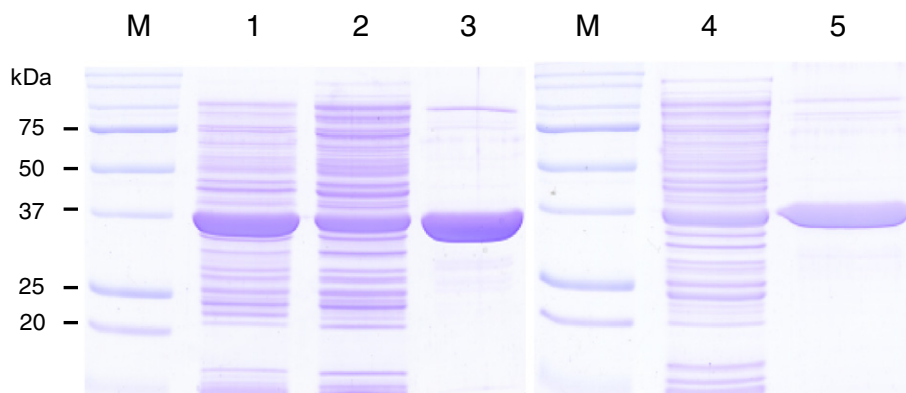


Fig. 4. Analysis of the expression of serine racemase genes from barley and rice in *E. coli* cells by SDS-polyacrylamide gel. Total cell lysate (lane 1), soluble protein (lane 2), purified enzyme following Ni-NTA column chromatography (CC) (lane 3) from *E. coli* cells harboring pHvSR, soluble protein (lane 4) and purified enzyme following Ni-NTA CC (lane 5) from *E. coli* cells harboring pOsSR. Gels were obtained using 15% SDS-PAGE with molecular mass marker series (lane M) for calibration, with detection by Coomassie Brilliant Blue R-250 staining.

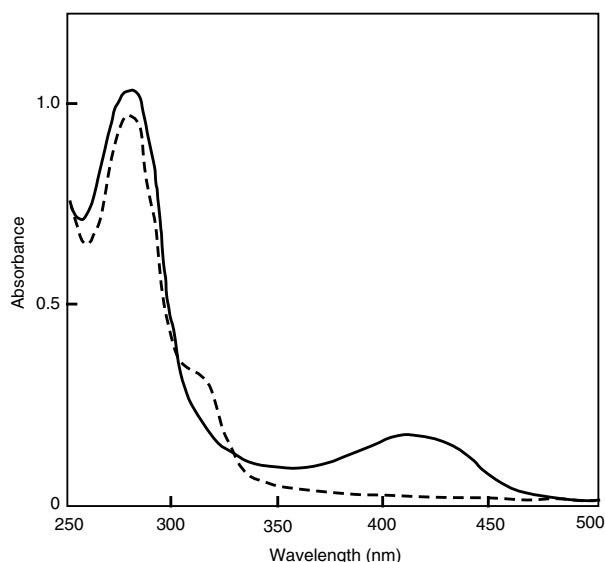


Fig. 5. Absorption spectra of purified HvSR before and after NaBH_4 incubation. The spectra of the purified HvSR (1 mg/ml) before and after dialyzing against 500 volumes of 20 mM Tris-HCl (pH 7.5) containing 1 mM DTT, 10 μM PLP, and 10 mM sodium borohydride are shown as solid and dashed lines, respectively.

2.4. Effects of divalent cations and ATP

The effects of divalent cations and ATP on **1b** racemization and dehydration activities were analyzed by adding 1 mM divalent ion to a reaction mixture at pH 8.5 (Fig. 6). Racemase activity was increased 2.9-, 3.8-, and 3.8-fold by addition of Mg^{2+} , Ca^{2+} , and Mn^{2+} , whereas EDTA, Fe^{2+} , or Ni^{2+} inhibited this activity. The addition of Mg^{2+} , Ca^{2+} , and Mn^{2+} also increased dehydratase activity 13-, 11-, and 13-fold, respectively, whereas Fe^{2+} showed only a 2.8-fold increase and Ni^{2+} inhibited the activity. HvSR was not activated by 1 mM ATP, and the activity with the combination of 1 mM Mg^{2+} and 1 mM ATP was slightly decreased compared to that in the presence of 1 mM Mg^{2+} .

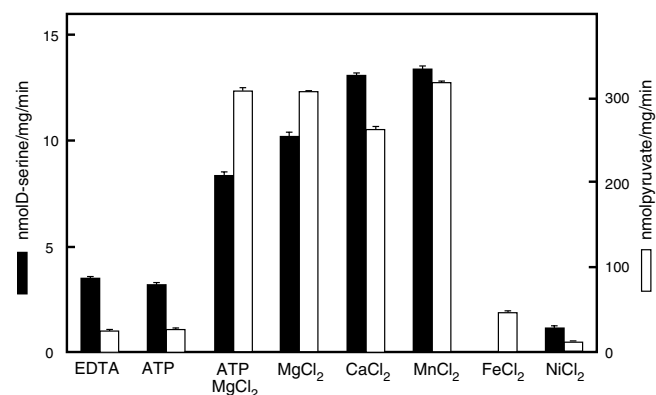


Fig. 6. Effects of divalent cations and ATP on racemase and dehydratase activities of barley serine racemase. Racemase (black columns) and dehydratase (white columns) activities of the purified recombinant enzyme were assayed in the presence of 1 mM EDTA, 1 mM metal chloride, or 1 mM ATP. The results represent the average \pm s.e. of three experiments.

2.5. Immunological properties

An immunological comparison among plant serine racemases was performed by Ouchterlony double diffusion (Fig. 7). Anti-HvSR serum formed one line not only with purified recombinant HvSR but also with purified recombinant OsSR and AtSR; however, the anti-AtSR serum also formed a precipitate with AtSR and no line was detected with HvSR and OsSR.

3. Discussion

We demonstrated that barley serine racemase, HvSR, is a bifunctional PLP-dependent enzyme that catalyzes not only racemization of **1a** and **1b** but also dehydration of **1a** and **1b** to **2**. The enzyme activity of HvSR and *A. thaliana* serine racemase, AtSR, are enhanced by addition of divalent cations such as Ca^{2+} , Mg^{2+} , and Mn^{2+} . Fe^{2+} inhibits racemase activity of HvSR, whereas that of AtSR

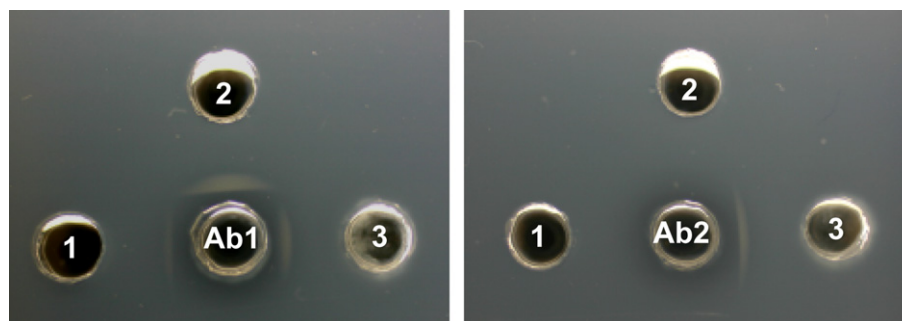


Fig. 7. Ouchterlony double diffusion of the purified recombinant plant serine racemases. Ab1, anti-HvSR serum; Ab2, anti-AtSR serum; 1, OsSR; 2, HvSR; 3, AtSR.

is slightly increased by addition of Fe^{2+} (Fujitani et al., 2006). While the combined effect of Mg^{2+} and ATP on mouse serine racemase leads to an increase in **1a** production, there was no synergy between Mg^{2+} and ATP on HvSR activity, as with AtSR. These results suggest that the protein structure interacting with ATP might be different between plant and mammalian serine racemases. The amino acid residues that interact with PLP (Phe61, Lys62, Asn89, Gly241, Ser315) are conserved in HvSR, with other plant and mammalian serine racemases, which belong to the fold type II group of PLP enzymes (Gallagher et al., 1998), except for the glycine-rich loop, which consists of a triglycine and a tetraglycine in plant and mammalian serine racemases, respectively. These results show that plant serine racemase represents a distinct group in the eukaryotic serine racemase family.

The amino acid sequence identities of HvSR with OsSR and AtSR (89% and 68%, respectively) and the phylogenetic tree suggest that plant serine racemases can be clustered into monocot and dicot types. To show the structural differences between serine racemases of monocot and dicot plants, we prepared anti-HvSR and anti-AtSR sera, and then purified recombinant rice serine racemase with both racemase and dehydratase activities: the latter was obtained by expression of an open reading frame of the putative rice serine racemase gene in DDBJ, EMBL, and GenBank data banks. The specific cross-reactivity of the anti-AtSR serum against AtSR by the Ouchterlony test supports the possibility that there may be monocot and dicot clusters in plant serine racemase based on a comparison of the three plant enzymes examined so far.

Recently, an aspartate racemase gene was cloned from a bivalve mollusk and expressed to characterize its enzyme properties (Abe et al., 2006). The aspartate racemase is a PLP-dependent enzyme and has both racemase and dehydratase activities. The amino acid sequence shows 38%, 39%, 32%, 30%, and 36 % identities to human, mouse, barley, rice, and *A. thaliana* serine racemases, respectively, and contains conserved PLP-interacting amino acid residues, corresponding to Phe61, Lys62, Asn89, Gly241, and Ser315 of *E. coli* threonine dehydratase. Interestingly, its enzyme activity was decreased by ATP but not by EDTA, whereas mammalian serine racemases was activated by

both divalent cations and ATP. In contrast to the eukaryotic racemases, plant serine racemase activity was increased by divalent cations but not by ATP. This suggests that the interaction of the enzyme with ATP and the divalent cation restricts the substrate specificity of eukaryotic racemases.

A novel mechanism of mammalian serine racemase has been proposed, whereby dehydratase activity of serine racemase regulates the intracellular **1a** level. This effect occurs especially in brain areas, since a D-amino acid oxidase, which degrades D-amino acids in mammalian cells, is restricted to the cerebellum and brainstem and a dehydratase activity-deficient serine racemase mutant accumulates **1a** (Foltyn et al., 2005). Several D-amino acids and D-amino acid-containing peptides have been identified in plants. However, both **1a** and D-alanine inhibit plant growth at low concentrations (Erikson et al., 2004), indicating that the D-amino acid level should be strictly regulated in plant cells. There is no previous report of a plant D-amino acid oxidase, and the serine racemase is the only known enzyme that degrades **1a** in plants. The specific activities of racemization and dehydration for **1b** were as follows: 14 and 228 nmol/min/mg with regard to HvSR, 10 and 214 nmol/min/mg with regard to OsSR, and 4.1 and 86 nmol/min/mg with regard to AtSR (Fujitani et al., 2006), respectively: the corresponding dehydratase activities were about 20-fold higher than racemase activity. This is in contrast to the dehydration/racemization ratio of about 0.7 for mouse serine racemase (De Miranda et al., 2002). These results thus suggest that the main physiological function of plant serine racemase might be to degrade, rather than produce **1a**.

4. Experimental

4.1. Plant material and growth conditions

The seeds of barley, *Hordeum vulgare* L., cv. Haruna nijo, were germinated and cultured in a hydroponic solution that consisted of 4 mM KNO_3 , 1 mM NaNO_3 , 1 mM NaH_2PO_4 , 1 mM MgSO_4 , 1 mM CaCl_2 , 45 μM Fe-citrate, 18 μM H_3BO_3 , 4.6 μM MnSO_4 , 1.5 μM ZnSO_4 , 1.5 μM CuSO_4 , and 1.5 μM Na_2MoO_4 . The pH of the cul-

ture solution was adjusted to 5.5 with NaOH. After 1 week of cultivation, the roots were harvested, frozen in liquid N₂, and stored at –80 °C.

4.2. Rapid amplification of cDNA ends PCR (RACE PCR)

Total RNA was isolated from the storage roots using the RNeasy Plant Mini Kit (Qiagen) and adaptor-ligated cDNA was synthesized from 1 µg of total RNA using the SMART RACE cDNA Amplification Kit (Clontech) according to the manufacturer's instructions. The primers for 3- and 5-RACE, 5'-GCCTTGAACCTTTGGAGCAAGTCCCTG-3' and 5'-TCCCTATTTTGCTGCTCTCATGCCAAG-3', respectively, were synthesized based on the conserved nucleotide sequences from truncated barley genes (accession nos. BJ450541 and CA023779) and PCR was performed on the cDNA with a synthesized primer and the universal primer mixture ("UPM") supplied in the kit. Fragments of about 280 and 800 bp amplified by 3- and 5-RACE PCR, respectively, were cloned into a pGEM-T vector (Promega) and subjected to nucleotide sequencing on both strands using a BigDye terminator v1.1 cycle sequencing kit (Applied Biosystems) with a series of synthetic primers.

4.3. Cloning of serine racemase gene

Based on the putative open reading frame of the barley serine racemase gene predicted by the RACE PCR experiment, primers were designed as follows: sense primer, 5'-GCCATATGGAAGCAGAGATGACGATG-3', which creates an *Nde*I site (underlined), and antisense primer, 5'-GACTCGAGTTTATACAAGGAATC-CCAT-3', which creates an *Xho*I site (underlined). The open reading frame was amplified by reverse-transcription PCR (RT-PCR) using the OneStep RT-PCR kit (Qiagen) with total RNA from the barley roots. The PCR product of 1026 bp was cloned into the pGEM-T vector and the *Nde*I- and *Xho*I-digested fragment of the plasmid was cut out and subcloned into an *Nde*I- and *Xho*I-digested pET20b(+) vector (Novagen), in which the endogenous stop codon of the serine racemase gene was replaced by a polyhistidine tag gene. The resulting plasmid, pHvSR, was transformed into *E. coli* BL21(DE3) pLysS cells.

The open reading frame of the rice serine racemase gene was amplified with total RNA from the rice, *Oryza sativa*, cv. Nipponbare, and the primers were designed based on the putative gene (AL606647), as follows: sense primer, 5'-CCCATATGGGGAGCAGAGGTGGAAGTGG-3', which creates an *Nde*I site (underlined), and antisense primer, 5'-CCCTCGAGACGTTTATAGAGAGACTCCC-3', which creates an *Xho*I site (underlined) by OneStep RT-PCR. The *Nde*I- and *Xho*I-digested fragment of the 1032 bp PCR product was subcloned into an *Nde*I- and *Xho*I-digested pCold I vector (Takara), in which a polyhistidine tag gene is fused upstream from the start codon. The resulting plasmid, pOsSR, was transformed into *E. coli* BL21 cells.

4.4. Expression and purification of serine racemase

E. coli cells harboring pHvSR were grown at 25 °C in Luria–Bertani (LB) medium containing 50 µg/ml of ampicillin. When the OD₆₀₀ became 0.5, isopropyl-β-D-thiogalactopyranoside (IPTG) was added to the culture at a final concentration of 0.5 mM. After cultivation at 25 °C for 6 h, the cells were harvested by centrifugation and frozen at –80 °C for at least 2 h. In the case of *E. coli* cells harboring pOsSR, the culture of 0.5 OD₆₀₀ was incubated at 15 °C for 30 min and cultured at 15 °C for 8 h with 0.5 mM IPTG. The frozen cell pellets were suspended in a BugBuster™ HT protein extraction reagent (Novagen) according to the manufacturer's instructions. The resulting recombinant protein was purified using an Ni-NTA CC (Qiagen) initially equilibrated in 20 mM Tris–HCl buffer (pH 7.9) containing 0.5 M NaCl and 5 mM imidazole (Buffer A). The column was next washed with Buffer A, followed by 60 mM imidazole in Buffer A, with the absorbed protein finally eluted with 1 M imidazole in Buffer A. The enzyme solution was collected, dialyzed against a 20 mM Tris–HCl buffer (pH 7.5) containing 1 mM DTT and 10 µM PLP, and concentrated by an Ultracent-30 (Tohso).

4.5. Enzyme and protein assays

The serine racemase and dehydratase activities were assayed by calculating the hydroxyperoxide production with amino acid oxidase–peroxidase–OPA and by calculating the pyruvate production with 2,4-dinitrophenylhydrazine, respectively, according to the method described previously (Fujitani et al., 2006). The **1b** solution used in the enzyme assay was pre-treated with D-amino acid oxidase and catalase to remove any contaminating **1a** according to De Miranda et al. (2002). L-Amino acid oxidase was used to remove contaminating **1b** in the **1a** solution. Protein concentrations were quantified according to Bradford (1976) with bovine serum albumin as the standard.

4.6. Antiserum preparation

The purified recombinant serine racemase of barley or *A. thaliana* was mixed with either Freund's complete adjuvant (for the first injection) or incomplete adjuvant (for following injections). The emulsified mixture containing protein (1 mg) was used to immunize a male Japanese white rabbit by subcutaneous injection followed by four subsequent injections, once every two weeks. Titration of the antiserum was performed by standard enzyme-linked immunosorbent assay (ELISA).

4.7. Computer analysis

Analysis and translation of the nucleotide sequences were performed using GENETYX-MAC (Software Development). The homology searches, alignments, and phylogenetic analyses of the amino acid sequences were

performed using BLAST (Altschul et al., 1990; Karlin and Altschul, 1990), CLUSTAL W (Tompson et al., 1994), and TreeView (Page, 1996) programs, respectively.

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