Metal Dependence of the Xylose Isomerase from *Piromyces* sp. E2 Explored by Activity Profiling and Protein Crystallography

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§Supporting Information

ABSTRACT: Xylose isomerase from *Piromyces* sp. E2 (PirXI) can be used to equip *Saccharomyces cerevisiae* with the capacity to ferment xylose to ethanol. The biochemical properties and structure of the enzyme have not been described even though its metal content, catalytic parameters, and expression level are critical for rapid xylose utilization. We have isolated the enzyme after high-level expression in *Escherichia coli*, analyzed the metal dependence of its catalytic properties, and determined 12 crystal structures in the presence of different metals, substrates, and substrate analogues. The activity assays revealed that various bivalent metals can activate PirXI for xylose isomerization. Among these metals, Mn$^{2+}$ is the most favorable for catalytic activity. Furthermore, the enzyme shows the highest affinity for Mn$^{2+}$, which was established by measuring the activation constants ($K_{a}$) for different metals. Metal analysis of the purified enzyme showed that in vivo the enzyme binds a mixture of metals that is determined by metal availability as well as affinity, indicating that the native metal composition can influence activity. The crystal structures show the presence of an active site similar to that of other xylose isomerases, with a D-xylose binding site containing two tryptophans and a catalytic histidine, as well as two metal binding sites that are formed by carboxylate groups of conserved aspartates and glutamates. The binding positions and conformations of the metal-coordinating residues varied slightly for different metals, which is hypothesized to contribute to the observed metal dependence of the isomerase activity.

Cost-effective production of second-generation bioethanol requires maximal utilization of sugars present in cellulosic biomass, because raw materials account for approximately one-third of the overall production cost. ¹ Besides D-glucose, the most abundant monosaccharide in lignocellulose and hemicellulose is D-xylose, and fermentation of xylose along with glucose would significantly increase the total ethanol yield. ² For such simultaneous fermentation of glucose and xylose, *Saccharomyces cerevisiae* would be a particularly attractive organism as it is an established ethanol producer that is not very vulnerable to inhibitors present in cellulose hydrolysates and shows a relatively high tolerance to extracellular ethanol. ³ However, natural strains of *S. cerevisiae* do not metabolize xylose, because of its inability to convert D-xylose to D-xylulose, an aldose to ketose isomerization reaction. Because D-xylulose can be metabolized, research has been devoted to engineer yeast variants that express a heterologous xylose isomerase for catalyzing this reaction. ⁴⁻⁶ Alternatively, xylose isomerization can be achieved by incorporating both a xylose reductase and a xylitol dehydrogenase, ⁷ although this is not preferred because it could cause a cellular cofactor imbalance. ⁸⁻⁹ Because xylose isomerase requires only metal cofactors, it offers the most attractive solution. Furthermore, the use of an isomerase rather than dehydrogenases prevents the formation of xylitol, which may emerge as an unwanted side product. ¹⁰

Xylose isomerase and related glucose isomerases have been studied for decades because of their application as glucose isomerases in the production of high-fructose corn syrups from starch hydrolysates. ¹¹ The use of the enzymes in ethanol-producing yeasts has been more recent. Isomerases from different bacterial sources, including *Escherichia coli*, *Actinoplanes missouriensis*, *Streptomyces rubiginosus*, *Bacillus subtilis*, and *Clostridium thermosulfurogenes*, have been tested, but functional expression in *S. cerevisiae* appeared to be problematic. ¹²⁻¹⁵ Xylose isomerase from *Thermus thermophilus* was functionally expressed in *S. cerevisiae*, but the enzyme from this thermophilic organism showed very low activity at 30 °C. ¹⁶ The first eukaryotic XI gene that could be expressed well in its active form in *S. cerevisiae* was obtained from the anaerobic fungus *Piromyces* sp. E2 (PirXI), and its expression in an *S.
cerevisiae enabled the fermentation of xylose to ethanol.\textsuperscript{5,17,18} Although later a few xylose isomerases from other sources were also expressed in \textit{S. cerevisiae},\textsuperscript{3,4,6,19,20} xylose-based growth rates and ethanol production levels do not exceed those found for the \textit{S. cerevisiae} expressing XI from \textit{Piromyces} sp. E2.\textsuperscript{18} A further improved \textit{S. cerevisiae} strain equipped with PirXI indeed showed high ethanol production yields, reaching almost the theoretical level, and the use of xylose isomerase appears to be preferable over the combination of a xylose reductase and a xylitol dehydrogenase allowing isomerization via xylitol.\textsuperscript{10,20}

In spite of these promising achievements, there is significant room to improve the anaerobic metabolism of xylose. Engineering of the strain by overexpressing the non-oxidative pentose phosphate pathway enzymes may enhance the growth rate.\textsuperscript{20,21} Evolutionary optimization can also improve the rate of xylose consumption.\textsuperscript{22,23} In such improved \textit{S. cerevisiae} strains, a very high expression level of PirXI is still required for growth on xylose, especially for anaerobic metabolism, and in such evolved strains, PirXI may be the major intracellular protein (unpublished data). This must be an energetic burden to the cells and also indicates that xylose isomerization still is the rate-limiting step in engineered strains that have lower expression levels of xylose isomerase.

XIs can be divided into two structural types, i.e., class I and class II enzymes. The major difference between the two classes is an N-terminal extension in class II xylose isomerases. From sequence analysis, the PirXI enzyme explored here appears to be a class II xylose isomerase. Currently, only a few class II XI structures are available,\textsuperscript{34} and these have no bound substrates, unlike structures for class I XIs that have been obtained with different ligands.

In this study, we describe the catalytic properties of PirXI expressed in \textit{E. coli}. We also examined the metal specificity of the enzyme by measuring catalytic activities and affinities with a range of different metals that might be present in the cytosol of \textit{S. cerevisiae} during the fermentation process. Furthermore, we report several structures of the class II XI from \textit{Piromyces}, determined by X-ray crystallography, with various combinations of metals (\textit{Mg}\textsuperscript{2+}, \textit{Mn}\textsuperscript{2+}, \textit{Co}\textsuperscript{2+}, \textit{Fe}\textsuperscript{2+}, \textit{Ca}\textsuperscript{2+}, \textit{Ni}\textsuperscript{2+}, and \textit{Cd}\textsuperscript{2+}) and substrate (xylose), product (xylulose), or inhibitor (xylitol or sorbitol). The results indicate that small differences in metal–protein interactions influence the position and reactivity of bound metals.

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**Figure 1.** Mechanism of conversion D-xylose to D-xylulose.\textsuperscript{25,26} Linearization of D-xylose catalyzed by His102/Asp105 is followed by movement of the catalytic metal from position M2a to position M2b, where it coordinates with O1 and O2 of the sugar. A proton is transferred from O2 to O1, and a hydride shifts from C2 to C1. D-Xylulose is formed by ring closure.
Expression of PirXI in *E. coli*. A synthetic *xylA* gene encoding XI from *Piromyces* sp. E2 (GenBank accession AJ249909) was obtained from GenScript USA Inc. (Piscataway, NJ). The sequence was codon-optimized for expression in *E. coli*. The *xylA* gene was cloned into a pBAD/myc-His-derived plasmid (Invitrogen). In this plasmid, the original NdeI sites were removed and the NcoI site was replaced with an Nhel site. The *xylA* gene was cloned using Ndel and HindIII sites with insertion of a stop codon at the end of the *xylA* coding sequence. The obtained pBAD-PirXI plasmid was transformed into *E. coli* Top10 (Invitrogen) or NEB 10-β (New England Biolabs). The sequence of the plasmid was verified by sequencing.

For expression of XI, *E. coli* Top10 cells or NEB 10-β cells harboring the pBAD-PirXI plasmid were cultivated in TB medium (12 g/L tryptone, 24 g/L yeast extract, 5 mM/L glycerol, 2.31 g/L KH2PO4 and 16.43 g/L K2HPO4·3H2O) supplemented with 50 μg mL−1 ampicillin and 0.2% (w/v) l-arabinine for 16 h at 37 °C. The cells were harvested by centrifugation, washed with 20 mM Tris-HCl (pH 8.5), and directly used for enzyme purification or stored at −20 °C until further use.

**Purification of XI Overexpressed in *E. coli***. The cells were disrupted by sonication, and cell debris was removed by centrifugation for 45 min at 31000 g and 4 °C. The supernatant was applied to a Q-sepharose Fast Flow resin (GE Healthcare) equilibrated with 20 mM Tris-HCl (pH 8.5) in a gravity flow column and incubated for 30 min at 4 °C in a rotating format. The column was washed with 15 column volumes of 20 mM Tris-HCl (pH 8.5). Then xylose isomerase was eluted from the column using 20 mM Tris-HCl (pH 8.5) containing 100 mM KCl. Fractions containing protein were pooled and desalted using an EconoPac 10-DG desalting column (Bio-Rad) pre-equilibrated with 10 mM MOPS (pH 7.0). Protein concentrations were determined using the theoretical extinction coefficient at 280 nm (ε280,xl = 73800 M−1 cm−1) calculated by the ProtParam tool (http://web.expasy.org/protparam/). The protein was stable when it was stored at 4 °C for several weeks. For long-term storage, the protein was frozen in liquid nitrogen and stored at −80 °C.

**Activity Assays**. Most assays were performed using the enzyme with a controlled metal composition. For this, we first prepared the apoenzyme by buffer exchange with 10 mM EDTA in 10 mM Tris-HCl (pH 8.0) using either dialysis or EconoPac 10-DG desalting columns (Bio-Rad). The samples were incubated with EDTA for 30 min. Subsequently, EDTA was removed by buffer exchange with 10 mM MOPS (pH 7.0). The activity of this apoXI assayed in the absence of added bivalent metal ions was <0.02 unit/mg.

In most cases, XI activities were measured by a coupled enzyme assay using D-sorbitol dehydrogenase (SDH). D-Xylose formed by XI activity is reduced by SDH to xylitol, which is monitored by following NADH oxidation spectrophotometrically at 340 nm. SDH was obtained from Roche Diagnostics GmbH (Mannheim, Germany). The standard reactions were performed at 30 °C, and the mixtures contained 20 mM MOPS (pH 7.0), 1 mM bivalent metal ions, 150 mM D-xylose, 0.15 mM NADH, and 1 unit/mL SDH. Reactions were initiated by adding 0.05—0.2 μM apoXI. The amount of XI was adjusted so that the ratio of SDH activity to XI activity would be at least 30. In the case of assays in the presence of Co2+, the amount of SDH was doubled to balance the inhibitory effect of Co2+ ions on SDH activity. The absorbance at 340 nm was followed using a spectrophotometer (Jasco) or a Synergy Mx microtiter plate reader (BioTek Instruments, Inc.). D-Xylose and D-fructose stocks (2 M) were made in Milli-Q water and stored at −20 °C. The measurements were performed in duplicate. One unit of XI activity is defined as the amount of enzyme catalyzing conversion of 1 μmol of D-xylose per minute under the assay conditions.

Kinetic parameters of XI for D-xylose were determined by measuring XI activity with varying D-xylose concentrations in the range of 0.04—1875 mM, depending on the metal cofactor used. For pH profiling of XI activities, different buffers (50 mM) were used to establish pH values: MES for pH 5.5 and 6.0 and MOPS for pH 6.5, 7.0, and 7.5. Activities of the native enzyme isolated from *E. coli* were measured in the presence of 1 mM MgCl2 using the spectrophotometric assay described above. For determining the apparent metal affinities (Kact values), the activity of apo-PirXI was measured in the presence of varying metal concentrations (1—4000 μM) using 150 mM xylose as the substrate for Mg2+ and Mn2+, 1375 mM xylose for Ca2+, and 15 mM xylose for Co2+. The concentration of the enzyme used was in the range of 0.05—0.2 μM for the reactions with Mg2+, Mn2+, and Co2+ and 0.5—1.0 μM for the reaction with Ca2+.

For measuring the activities of PirXI with metal cofactors that are not compatible with the SDH-coupled assay, activities were examined by measuring the formation of D-xylulose from D-xylose by high-performance liquid chromatography (HPLC). Reaction mixtures containing the substrate, PirXI, and 1 mM metal ions in MOPS (pH 7.0) were incubated at 30 °C while being shaken. Reactions were stopped by acidification with HCl and addition of acetonitrile to a final concentration of 80%. Samples were then separated on an XBridge BEH Amide column (4.6 mm × 250 mm, pore size of 130 Å, particle size of 3.5 μm, Waters). Separation conditions were isocratic with 80% acetonitrile and 20% 20 mM MOPS (pH 7.0) and a flow rate of 1 mL/min at 80 °C. The ultraviolet (UV) absorbance at 210 nm was measured for D-xylulose quantification. For measuring XI activity with Fe3+, oxygen-limited reaction mixtures were prepared by flushing the reaction components and the mixture with argon. The reaction mixtures were incubated in a 30 °C water bath. During the reaction, the argon was flushed over the surface of the reaction mixture, and for sampling, a syringe was used. The activity of PirXI on glucose was also measured using the HPLC assay as described above. The activities with different concentrations of glucose ranging from 0.1 to 1.5 M were measured in the presence of 1 mM MgCl2 as the metal cofactor. A refractive index detector was employed for D-fructose quantification.

**Metal Analysis**. XI purified from *E. coli* was lyophilized and analyzed for the presence of calcium, cobalt, copper, iron, magnesium, manganese, and zinc using inductively coupled plasma optical emission atomic spectroscopy on an Optima 7000DV ICP-OES apparatus (PerkinElmer). The measurements were performed in duplicate. Before lyophilization, a molybdenum standard (molybdenum ICP standard Certipur, Merck Millipore) was added to the sample to a final concentration of 5.0 μg/mL.

**Crystallization and Structure Determination**. Initial vapor-diffusion crystallization experiments were performed using a Mosquito crystallization robot (TTT Labtech). In a typical experiment, 0.1 μL of the screening solution was added...
### Table 1. Data Collection and Refinement Statistics

<table>
<thead>
<tr>
<th>Sample</th>
<th>Cryoprotectant</th>
<th>Unit Cell Dimensions</th>
<th>R/Refl (%)</th>
<th>Resolution (Å)</th>
<th>Completeness (%)</th>
<th>I/σ(I)</th>
<th>Redundancy</th>
<th>R/Rfree (%)</th>
<th>active site occupancy</th>
</tr>
</thead>
<tbody>
<tr>
<td>native</td>
<td>20% glycerol</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>12.1/14.8</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>20.6</td>
<td>3.9</td>
<td>115.4, 90.0, 117.0</td>
<td>4 (Ca2+,Mg2+,Fe2+)</td>
</tr>
<tr>
<td>apo</td>
<td>20% glycerol</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>18.4/21.0</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>9.1</td>
<td>4.0</td>
<td>175.1/185</td>
<td>0.4, 0.35, 0.25</td>
</tr>
<tr>
<td>Mg−glycerol</td>
<td>0.5 M xylitol</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>15.5/17.6</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>8.4</td>
<td>4.0</td>
<td>14.0/16.1</td>
<td>4 Mg2+,Mg2+,Fe2+</td>
</tr>
<tr>
<td>Mg−xylitol</td>
<td>2.2 M xylitol</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>15.5/17.6</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>14.0</td>
<td>4.0</td>
<td>14.0/16.1</td>
<td>4 Mg2+,Mg2+,Fe2+</td>
</tr>
<tr>
<td>Mg−xylose</td>
<td>2.2 M xylose</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>15.5/17.6</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>6.1</td>
<td>4.0</td>
<td>14.0/16.1</td>
<td>4 Mg2+,Mg2+,Fe2+</td>
</tr>
<tr>
<td>Ca−xylose</td>
<td>2.2 M xylose</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>15.5/17.6</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>9.3</td>
<td>4.0</td>
<td>14.0/16.1</td>
<td>4 Mg2+,Mg2+,Fe2+</td>
</tr>
<tr>
<td>Mn−xylose</td>
<td>2.2 M xylose</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>15.5/17.6</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>9.3</td>
<td>4.0</td>
<td>14.0/16.1</td>
<td>4 Mg2+,Mg2+,Fe2+</td>
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<tr>
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<td>2.2 M sorbitol</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>15.5/17.6</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>9.3</td>
<td>4.0</td>
<td>14.0/16.1</td>
<td>4 Mg2+,Mg2+,Fe2+</td>
</tr>
<tr>
<td>Fe−glycerol</td>
<td>2.2 M xylose</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>15.5/17.6</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>9.3</td>
<td>4.0</td>
<td>14.0/16.1</td>
<td>4 Mg2+,Mg2+,Fe2+</td>
</tr>
<tr>
<td>Ca−xylose</td>
<td>2.2 M xylose</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>15.5/17.6</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>9.3</td>
<td>4.0</td>
<td>14.0/16.1</td>
<td>4 Mg2+,Mg2+,Fe2+</td>
</tr>
<tr>
<td>Ni−xylose</td>
<td>2.2 M xylose</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>15.5/17.6</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>9.3</td>
<td>4.0</td>
<td>14.0/16.1</td>
<td>4 Mg2+,Mg2+,Fe2+</td>
</tr>
<tr>
<td>Cd−xylose</td>
<td>2.2 M xylose</td>
<td>a = 78.7, b = 79.4, c = 92.1</td>
<td>15.5/17.6</td>
<td>1.80</td>
<td>92.3 (75.6)</td>
<td>9.3</td>
<td>4.0</td>
<td>14.0/16.1</td>
<td>4 Mg2+,Mg2+,Fe2+</td>
</tr>
</tbody>
</table>

*Values in parentheses are for the highest-resolution shell.*
to 0.1 μL of the protein solution on a 96-well MRC2 plate (Molecular Dimensions); reservoir wells contained 50 μL of the screening solution. The screening solutions used for the experiments were PACT and JCSG+ (Qiagen Systems). Crystallization conditions were optimized using hanging-drop setups with 13–15% (w/v) PEG 3350 and 0.1 M ammonium sulfate in 0.1 M HEPES (pH 7.0) as a precipitant, and 1 μL of the reservoir solution at 295 K. Native PirXI crystals belong to space group C2, with a solvent content of 45%. Apo crystals were grown automatically in triethanolamine (pH 7.6) and 1.295 K. Native PirXI crystals belong to space group Xylulose was available only as a 0.5 M solution.

Single cryo-cooled crystals were mounted on an in-house MarDTB Microstar rotating-anode generator equipped with HeliosMX Mirrors. Intensity data were processed using iMosflm38 and scaled using Aimless.39 Phases were obtained by molecular replacement using Phaser.40 The coordinates of three xylose isomerase structures [Protein Data Bank (PDB) entries 1A0C, 1A0D, and 1A0E], which have sequences that are ∼50% identical with that of PirXI, were used to construct a composite search model. The PirXI structure was built using ARP/WARP41 and Coot42 and refined using Refmac5.43 Water molecules were placed automatically in Fo − Fc difference Fourier maps at a 3σ cutoff level and validated to ensure correct coordination geometries using Coot. The first residues from the four XI subunits were not visible in the electron density and therefore not included in the final model. In the binary and ternary complexes, 2Fo − Fc and Fo − Fc maps showed extra density in the active site. To confirm the metal ions, we calculated anomalous difference Fourier maps, with data collected at a wavelength of 1.54 Å (1.3 anomalous electrons for Ca, 2.8 for Mn, 3.2 for Fe, 3.7 for Co, 0.5 for Ni, and 4.7 for Cd), using phases obtained from the model without any metal ions. Occupancies of the metal ions were calculated with Phenix.44 Relevant statistics of data collection and model refinement are listed in Table 1. The stereochemical quality of the model was assessed with MolProbity.45 Interface areas between the subunits were calculated with PDBProSA.46 Figures were prepared with PyMOL (http://www.pymol.org) and ESPript.47 Atomic coordinates and experimental structure factor amplitudes have been deposited in the PDB (Table 1).

**Structure-Based Sequence Alignment.** A structure-based alignment of class II xylose isomerases from *Piromyces* sp. E2, *Bacteroides thetaiotaomicron* (PDB entry 4XK5), *Thermaanaerobacterium thermosulfurigenes* (PDB entry 1AOC), *Thermotoga neapolitana* (PDB entry 1A0E), and *Bacillus stearothermophilus* (PDB entry 1A0D) and a class I xylose isomerase from *St. rubiginosus* (PDB entry 4W4Q)38 was made with Promals3D.49 The alignment figure was created with ESPript.50

**RESULTS**

**Expression of PirXI in E. coli and Enzyme Isolation.** To overexpress the *Piromyces* sp. E2 xylose isomerase (PirXI) in *E. coli*, a codon-optimized synthetic *xylA* gene was cloned into the pBAD vector and transformed into *E. coli* Top10. Overnight cultivation at 37 °C with induction of PirXI by 0.2% arabinose resulted in high levels of soluble PirXI. Sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE) analysis showed that PirXI is the predominant protein in cell lysates, accounting for >50% of the total soluble protein. This allowed purification by a single step of ion-exchange chromatography, which after desalting routinely yielded 300–500 mg of purified xylose isomerase per liter of culture. From this material, the apoenzyme was prepared by treatment with EDTA to study the metal dependence of the isomerase activity. The purified enzyme was stored at −80 °C for use in further biochemical experiments.

SDS–PAGE showed that the size of the protein monomer is around 50 kDa, which is in good agreement with the calculated size of the protein, 49.5 kDa. Analytical size exclusion chromatography revealed a molecular weight of ~200 kDa for the native PirXI, showing that the enzyme is a tetramer.

**Metal Dependence of PirXI Activity.** Because xylose and glucose isomerases require bivalent metals for activity,51 and the type of metal that is bound may influence the activity,52 we

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**Table 2. Metal Dependence of PirXI Activity**

<table>
<thead>
<tr>
<th>substrate</th>
<th>cation</th>
<th>V (200 mM) (s⁻¹)</th>
<th>kcat (s⁻¹)</th>
<th>K₅₀ (mM)</th>
<th>kcat/K₅₀ (s⁻¹ M⁻¹)</th>
<th>K₅₀ (μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>xylose</td>
<td>Mg²⁺</td>
<td>1.95 ± 0.12</td>
<td>2.0 ± 0.1</td>
<td>7.5 ± 1.2</td>
<td>270</td>
<td>98.9 ± 4.3</td>
</tr>
<tr>
<td>xylose</td>
<td>Mn²⁺</td>
<td>4.4 ± 0.42</td>
<td>4.5 ± 0.3</td>
<td>4.3 ± 0.1</td>
<td>1100</td>
<td>1.45 ± 0.28</td>
</tr>
<tr>
<td>xylose</td>
<td>Ca²⁺</td>
<td>0.075 ± 0.017</td>
<td>0.8 ± 0.1</td>
<td>1930 ± 70</td>
<td>0.2</td>
<td>159 ± 10</td>
</tr>
<tr>
<td>xylose</td>
<td>Al³⁺</td>
<td>1.0 ± 0.14</td>
<td>1.0 ± 0.1</td>
<td>0.3 ± 0.04</td>
<td>3300</td>
<td>6.5 ± 1.21</td>
</tr>
<tr>
<td>glucose</td>
<td>Mg²⁺</td>
<td>0.032 ± 0.005</td>
<td>0.1 ± 0.01</td>
<td>430 ± 10</td>
<td>0.2</td>
<td>–</td>
</tr>
<tr>
<td>xylose</td>
<td>Zn²⁺</td>
<td>1.1 ± 0.1</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>xylose</td>
<td>Fe²⁺</td>
<td>1.6 ± 0.2</td>
<td>–</td>
<td>–</td>
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<td>–</td>
</tr>
<tr>
<td>xylose</td>
<td>Cd²⁺</td>
<td>0.2 ± 0.01</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>xylose</td>
<td>Ni²⁺</td>
<td>nd</td>
<td>–</td>
<td>–</td>
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</tr>
<tr>
<td>xylose</td>
<td>Al³⁺</td>
<td>nd</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

*The values represent averages and the mean deviation calculated from two or three replicates. Initial rates at 200 mM substrate were calculated from steady state parameters. K₅₀ is defined as the metal concentration at which the enzyme activity is half the V₅₀.*

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examined the effect of different metal ions on the activity of PirXI. Kinetic parameters of apo-PirXI in the presence of Mg2+, Mn2+, Co2+, or Ca2+ were measured using the SDH-coupled assay as described in Materials and Methods.

PirXI showed good activity in the presence of Mg2+, Co2+, and Mn2+ (Table 2). Among these metals, the highest activity (kcat) of the enzyme was obtained with Mn2+, followed by Mg2+ and Co2+. The substrate affinity with xylose (Km) on the other hand, was best in the presence of Co2+, which gave a KmA of 0.3 mM and also the highest catalytic efficiency (kcat/Km). In the presence of Mn2+ or Mg2+, the KmA of the enzyme was 15–20-fold higher. This metal preference of PirXI is similar to what was described for other class II XIIs, which were reported to be most active with Mn2+, whereas the class I enzymes are most active with Mg2+. When Ca2+ was added as the metal cofactor of PirXI, the enzyme showed activity, although it was lower than with the other metals mentioned above. A striking observation was the extremely low xylose affinity (KmA ~ 2 M) in the presence of Ca2+, resulting in a poor catalytic efficiency (kcat/KmA = 0.2 s−1 M−1). The low substrate affinity suggests that the Ca2+-containing enzyme will provide a negligible contribution to in vivo conversion of xylose by PirXI.

Further assays for determining the metal dependence of PirXI activity were performed using an HPLC assay for measuring xylulose formation because the SDH-coupled assay is not compatible with the presence of Zn2+, Fe2+, Cd2+, Ni2+, and Al3+ as these metals inhibit SDH activity. Furthermore, in the case of Fe2+-dependent activity, the reactions were performed under reduced oxygen conditions to prevent oxidation of Fe2+ to Fe3+. In these experiments, we measured initial conversion rates (units per milligram) of xylose that was added to a final concentration of 200 mM. PirXI showed activity in d-xylene isomerization in the presence of Zn2+, Fe2+, Cd2+, and Mn2+. The rate of xylose conversion with Zn2+ was almost half of that measured with Mg2+. Furthermore, Fe2+ gave moderate activity compared to activities with other metals. Unexpectedly, the enzyme also showed some activity in the presence of Cd2+, which was reported to be an inhibitor of another xylose isomerase.18 However, the activity (0.2 unit/mg) was very low. No detectable amount of xylulose (<0.5 μM) was found with Ni2+ or Al3+ as the metal cofactor upon measurement after incubation of the reaction mixture for 2 h.

Metal Affinity of PirXI. The results described above show that PirXI can be activated by a range of different bivalent metal ions. Obviously, the activity in vivo will also be determined by metal affinity and metal availability in the cytoplasmic environment. Therefore, we investigated the activity of PirXI on xylose in the presence of different concentrations of divalent metals and determined half-saturation constants for activation of the enzyme [KmA values (Table 2)].

Of the metals tested (Mg2+, Mn2+, Co2+, and Ca2+), the highest affinity was found with Mn2+. The enzyme also showed a high affinity for cobalt, but the affinities for Mg2+ and Ca2+ were 70- and 100-fold lower compared to that of Mn2+, respectively.

Analysis by inductively coupled plasma atomic emission spectroscopy showed that the main divalent metal in native PirXI isolated from E. coli is Mg2+, which accounted for approximately half of the total metal content (Table 3). Ca2+ and Fe2+ were also detected, whereas the enzyme contained only a few percent (moles per mole) of Mn2+ and Zn2+. This indicates that Mg2+ would strongly influence the in vivo enzyme activity in E. coli, while Mn2+ ions, which are most beneficial for activity, would have little influence. In agreement with this, the native enzyme isolated from E. coli showed activity similar to that of the enzyme reconstituted with Mg2+ or even slightly lower activity, probably due to the effect of Ca2+ and Fe2+. The metal composition and the activity of the enzyme are slightly different from one batch of isolated enzyme to another (WT1–WT3). The data also suggest that intracellular metal availability influences the metal composition of the enzyme (Mg > Ca > Mn > Co) as much as the relative affinities (Mn > Co > Mg > Ca). Accordingly, in vivo, the enzyme may have suboptimal activity because of the level and type of metal binding. Recently, Verhoeven et al.35 showed that the increase in the intracellular Mn2+ level improved the growth of S. cerevisiae on xylose by enhancing PirXI activity.

pH Dependence of PirXI Activity. The cytosolic pH of S. cerevisiae during the fermentation process is <7.0, possibly because of the influence of acidic lignocellulose hydrolysates.52,53 We measured the activity of PirXI at different pH values ranging from 5.5 to 7.5 in the presence of Mg2+, Mn2+, Ca2+, or Co2+ (Figure 2). Because of the high KmA of the enzyme in the presence of Ca2+, the activity was measured at a higher xylose concentration, 1 M instead of 150 mM. With Mg2+, Mn2+, and Ca2+, the results show a similar broad pH profile, with the highest activity at pH 7.5, and a significantly reduced activity at pH 6.5. The lower activity at low pH was more pronounced for Mg2+, the quantitatively most important cation in the non-reconstituted enzyme. The activity of the Co2+-reconstituted PirXI was much less dependent on pH in the range of 5.5–7.5.

Crystal Structure of PirXI. PirXI isolated from E. coli readily crystallized from PEG 3350 at pH 7. The structure was determined by using X-ray diffraction and molecular replacement and was refined against 1.8 Å resolution diffraction data to an R factor of 0.116 (Rfree = 0.148) with good stereochemistry (Table 1 and Figure 3A). The P1 unit cell contains four monomers that form a 92 Å × 77 Å × 77 Å tetramer, having noncrystallographic 222 symmetry. The four subunits have root-mean-square deviation (rmsd) values of 0.11–0.16 Å on Cα atoms. Each PirXI subunit comprises two structural domains, a larger catalytic domain (residues 1–377) with a distorted (β/α)8-barrel (TIM-barrel) fold and a C-terminal domain (residues 378–437) forming an extended tail containing three α-helices. With other class II xylose isomerases with PDB entries of 4XKM (B. thetaiotaomicron),34 1AOC (T. thermosulfurigenes), 1AOD (B. stearothermophilus), and 1AOF (Th. neapolitana), rmsd values are 0.4 Å on 435 Ca atoms, 1.2 Å on 430 Ca atoms, 1.1 Å on 425 Ca atoms, and 1.2 Å on 422 Ca atoms.
respectively, while with the well-studied St. rubiginosus class I enzyme, the rmsd is 2.0 Å on 352 Cα atoms and 27% sequence identity. Major differences between the class I St. rubiginosus XI and the class II PirXI are the presence in the latter enzyme of an N-terminal extension of ∼33 residues, an N-terminal addition of α-helix 1, and a larger loop before α-helix 2. Furthermore, the C-terminal regions deviate the most with low identity, except for the ultimate α-helix 12 (Figure S1).

A pair of subunits forms the “yin-yang” dimer with rotational symmetry by contacts of the catalytic domain of subunit A interacting with the C-terminal tail of subunit B, while the core of subunit B is interacting with the C-terminal tail of subunit A (Figure 3B). The other dimer pairs are called the “butterfly” dimer (Figure 3C) and the “diagonal” dimer (Figure 3D). The tetramer has a surface area of 53920 Å², of which 36840 Å² is buried. Each subunit has a total solvent accessible surface area of 21100 Å², of which 8110 Å² is buried. The interactions in the yin-yang and diagonal dimers are mainly present between the N- and C-terminal tails of the subunits, while the butterfly dimer has interactions between the cores of the subunits.

The sizes of the interface areas are similar in other class II xylose isomerases for which the structures have been determined (PDB entries 4XKM, 1A0C, 1A0D, and 1A0E) (Table 4). However, the xylose isomerase from St. rubiginosus, the most studied class I XI in the PDB, has a larger interface for the yin-yang dimer (4670 Å²) and a smaller interface for the diagonal dimer (1460 Å²) as it is missing the N-terminal extension of the class II enzymes. Class II enzymes, having larger dimer interfaces and interactions with the other subunit, should be regarded as tetramers unlike class I enzymes, which are usually described as a dimer of dimers.34

**Active Site Structure.** The active sites of the PirXI subunits are located on the C-terminal end of the (β/α)₈-barrel of the core domain. The active sites are 36 Å apart in yin-yang, 31 Å apart in butterfly, and 39 Å apart in diagonal dimers. Residues from the loops following the β-strands shape the active site. The active site pocket is lined by the side chain indole groups from Trp50 and Trp189 that are situated 8.8 Å from each other forming a cassette for sugar binding (Figure 4A,B). Other hydrophobic residues surrounding the substrate binding site are Phe61 from another subunit of the butterfly dimer, Phe146 and Trp140.

In the crystals obtained with native PirXI isolated from E. coli, only the position of the structural metal (M1) in the active site is occupied. Residues Glu233, Glu269, Asp297, and Asp340 respective, while with the well-studied St. rubiginosus class I enzyme, the rmsd is 2.0 Å on 352 Cα atoms and 27% sequence identity. Major differences between the class I St. rubiginosus XI and the class II PirXI are the presence in the latter enzyme of an N-terminal extension of ∼33 residues, an N-terminal addition of α-helix 1, and a larger loop before α-helix 2. Furthermore, the C-terminal regions deviate the most with low identity, except for the ultimate α-helix 12 (Figure S1).

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In the crystals obtained with native PirXI isolated from E. coli, only the position of the structural metal (M1) in the active site is occupied. Residues Glu233, Glu269, Asp297, and Asp340
bind M1 with octahedral geometry (Figure 4A). The nature of the metal ion was unclear from the structure as a mixture of different metals may bind. The observed metal–donor distances were 2.1–2.2 Å. The optimal distance for Ca²⁺ is 2.36 Å, for Mg²⁺ 2.26 Å, and for Fe²⁺ 2.01 Å to monodentate Asp or Glu.54 An anomalous difference map showed a peak at the M1 position indicating a bound Ca²⁺ or Fe²⁺ ion. Reﬁning with a Ca²⁺ ion with full occupancy showed residual Fo − Fc density and distances that were too short, while reﬁning with 100% Fe²⁺ showed deﬁcit density and distances that were too long. Therefore, M1 was reﬁned to occupancies of 0.4, 0.35, and 0.25 for Ca, Mg, and Fe, respectively, which gave a ﬂat Fo − Fc map after reﬁnement. The ambiguous metal ion content has also been observed in native crystals of St. rubiginosus with 0.6 equiv of Mn, <0.6 equiv of Mg, and <0.1 equiv of Co per monomer.55 In this PirXI structure, a glycerol molecule from the cryoprotectant is bound in the active site to the metal ion with a Me−O₂ distance of 2.3 Å and a Me−O₃ distance of 2.4 Å. Glycerol, acting as a sugar mimic, is often found in active sites of carbohydrate-converting enzymes.56

The crystal structure obtained with EDTA-treated XI is highly similar to the native structure with an rmsd of 0.1 Å. As expected, this apoenzyme does not contain any metals in the active site, but a glycerol molecule is bound as in the native enzyme. A water molecule is located at position M1; consequently, O3 of the glycerol has shifted 1.4 Å toward His272 (Figure 4B).

To determine if and how other metals bind to PirXI, crystal soaks were performed. Several structures of binary and ternary complexes of PirXI with metals, substrates, and inhibitors were determined (Table 1 and Table S1).

Mg, Xylose, and Xylitol Binding. In electron density maps of crystals of the apoenzyme soaked with magnesium and glycerol or with magnesium and the competitive inhibitor (or substrate analogue) xylitol,28 only a single Mg²⁺ ion at the M1 site was observed (Figure 5A,B). There was no 2Fo − Fc density present for the Mg²⁺ ion at position M2, despite the presence of Mg²⁺ in the soak solution at 10 mM (Figure S2). Glycerol was bound as in the native PirXI crystals. The position of xylitol was overlapping with that of the glycerol molecule in the Mg−glycerol structure, but the larger xylitol molecule bound deeper in the active site toward the putative M2 position. Xylitol was bound with its O₁ atom bound to NE₂ of His272 and a water molecule, O₂ to Glu233, Glu269, Asp340, and M₁ (Mg²⁺), O₃ to Asp340 and a water molecule, O₄ to Glu233 (bidentate), Asp297, Asp340, and M₁, and O₅ to NE₂ of His102 and a water molecule (Figure 5B). His102 is likely involved in pyranose ring opening.26,57

Only when a soak was performed with magnesium and the substrate D-xylose was the second metal site (M2) fully occupied with elongated (elliptical) electron density, showing two distinct positions for the metal, with approximately 50/50 occupancy, at distances of 3.8 Å (M2b) and 5.5 Å (M2a) from M1 (Figure 5C). It has been elucidated by structural studies, including neutron diﬀraction, that the hydride shift of the isomerization reaction is mediated by the metal at M2.25,26,58,59

In the Mg−xylose structure, the substrate is in its open form in the same conformation as xylitol, indicating ring opening has been completed. At the current resolution of the data sets (1.9 Å), it is not possible to distinguish between the linear forms of the substrate and product by electron density, and probably a mixture of both is bound. The open chain forms are similar in shape, and differences exist at only C1 and C2, having sp² or sp³
hybridization. Because all soaks were performed with a high concentration (2.2 M) of substrate and the rate-limiting step in the catalytic cycle was proposed to be the hydride-shift reaction, we have modeled the substrate used in the soak. The xylose stacks on the ring system of Trp189, which probably ensures a flat conformation of the substrate (Figure 5C). The M2b metal has interactions with O1 and O2 of the xylose, NE2 of His272, Glu269 (bidentate), and a water molecule. The metal at M2b is in the catalytically more favorable position as the interactions with O1 and O2 likely stabilize the transition state while the hydride is shifting from C2 to C1. These interactions with O1 and O2 of xylose are absent when the metal is at position M2a. It was therefore suggested that the metal moves from position M2a to position M2b during the reaction. Mg at M2a is 2.0 Å from M2b and has interactions with NE2 of His272, OE2 of Glu269, OD1 of Asp310, Asp308 (bidentate) in one of its conformations, and the water molecule. In another conformation, Asp308 and O1 of the xylose have interactions with NZ of Lys235, which has moved closer into the active site compared to its position in the Mg–xylitol structure (Figure 5C). Glu238, having a H-bond with the backbone amide of Asp308, also has achieved a double conformation. Hence, its structural neighbor, the side chain of the protein environment are shown as dashed lines. The difference electron density, $F_o - F_c$ contoured at the 2σ level for the sugars is shown. 

Figure 6. Active sites of (A) Cd–linear XLS–XI and (B) Cd–cyclic XLS–XI. The residues involved in the metal binding sites are shown as green sticks. His102 is colored salmon. The xylose molecules are colored yellow. Cd ions are depicted as light orange spheres, and interactions between the metal ions and the hydroxyl groups of xylose or the protein environment are shown as dashed lines. The crystal structure of XI–Cd with the xylose in a linear form is very similar to the structure of Mg–xylitol structure (Figure 5C). The residues involved in the metal binding sites are shown as green sticks. His102 is colored salmon. The xylose molecule is colored yellow. The residues and sorbitol from 2XIN are colored gray. Mn2+ ions are shown as purple spheres and Co2+ ions as pink spheres. 

Figure 7. Overlay of the active sites of Mn–SOR–XI and XI from A. missouriensis complexed with Co2+ and sorbitol (PDB entry 2XIN). The residues involved in the metal binding sites are shown as green sticks. His102 is colored salmon. The sorbitol molecule is colored yellow. The residues and sorbitol from 2XIN are colored gray. Mn2+ ions are shown as purple spheres and Co2+ ions as pink spheres. Interactions between the metal ions and hydroxyls of the sorbitol are shown as dashed lines.

In the Mg–xylitol complex of PirXI, bound substrates are in a linear form (XLS) in subunit A and subunit B, while in subunit C and subunit D of the yin-yang dimer, the substrates are in the pyranose ring form (Figure 6A,B). This mixture of open and closed structures was observed for only the PirXI–Cd structure. For PirXI structures soaked with other metals, the electron density map showed only the linear form of the substrate in the active site (Figure 5B–D). The active site structure of PirXI–Cd with the xylose in a linear form is very similar to the structure of Mg–xylitol. Cd at M2 is observed at two distinct positions with a 1.4 Å distance between M2a and M2b.

The electron density map in the active centers of subunits C and D shows α-D-xylopyranose (XYS) that is coordinated to M1 with O3 and O4, whereas endocyclic O5 is hydrogen bonded to His102. O4 of α-D-xylopyranose is also coordinated to M2b at a distance of 2.7 Å (Figure 6B).

The crystal structure of XI–Mn soaked with sorbitol, a C6 sugar analogue with the same stereo configuration as α-D-glucose, shows two singly occupied metal binding sites (Figure 7). The binding of the sorbitol is quite different from the binding of xylitol in the Mg–xylitol structure. O1 of sorbitol interacts with His102 (2.6 Å) and a water molecule, and O2 interacts with Glu233 (bidentate), Asp297, and M1-Mn2+. O3 is ligated by Glu233, Asp340, and M1, and O4 is ligated by Asp340 (bidentate) and Ne of Trp50. O5 and O6 have interactions with water molecules. Interactions with M2 occur via water molecules. The binding mode of sorbitol is also quite different from that in class I XI from A. missouriensis complexed with Co2+ and sorbitol (Figure 7) and from St. rubiginosus complexed with Ni2+ and sorbitol. In class II enzymes, the bulkier Trp140 replaces a Met (Figure 5I) in the rear side of the active site. Therefore, steric hindrance was expected in the active site of class II XI as the C6 atom of sorbitol would be too close to Trp140. In PirXI, the sorbitol molecule has shifted by 2.3 Å toward the entrance of the active site. Assuming that its C6 analogue binds in a similar manner, we found that glucose would bind to the active site in a less productive mode.
**Biochemistry**

**XI–Ni, XI–Fe, and XI–Ca.** The XI–Fe structure contains two iron atoms in the active site 1.9–2.2 Å from the surrounding ligands (Figure S3A). No glycerol molecules could be observed in the electron density, which could be caused by the lower resolution of the data set. The M2 Fe$^{2+}$ is positioned at the M2b location 2.4 Å from His272. The Fe$^{2+}$ ions are positioned 3.8 Å from each other.

A soak with xylene of a Ni$^{2+}$-grown XI crystal reveals xylene in its linear form in the active site (Figure S3B). Its position superimposes well on the Mg$^{2+}$-xylene structure. The Ni$^{2+}$ cation at position M2 has two distinct locations (M2a and M2b), as in the Mg$^{2+}$-xylene structure. However, Asp308 does not have the double conformation as observed in the Mg$^{2+}$-xylene structure.

A soak with Ca$^{2+}$ resulted in both positions M1 and M2 being occupied (Figure 8A). M2 is bound in a distorted octahedral geometry by His272 at 3.4 Å, Glu269 at 2.3 Å, Asp308 at 2.2 Å, Asp310 at 2.4 Å, and two water molecules. To the best of our knowledge in Class I XIs, the M1 site was never occupied by a Ca$^{2+}$ ion. At M2, other class I XIs bound Ca$^{2+}$ at the M2a position, similar to what we observed in the PirXI–Ca$^{2+}$–xylene structure, but the equivalent residue of Asp308 binds the M2 in a bidentate manner in other enzymes. The large size of M2–Ca$^{2+}$ shifts Asp308 and Asp310 slightly out and the O1 position of the linear xylene by 1.5 Å toward Phe61′. This leaves the substrate in an unfavorable position for isomerization.

**XI–Mn, XI–Co, and XI Substrate/Product.** The PirXI structure with two manganese ions and the substrate xylene well represents the most active form of the enzyme (Figure 8A). The Mn$^{2+}$ at position M1 has its usual octahedral coordination, while M2 has full occupancy at the catalytic M2b position (no occupation of the M2a site). The M1 and M2b metal binding sites are 4.1 Å distance from each other. M2b is coordinated by His272 at 2.4 Å, Glu269 at 2.3 and 2.9 Å, Asp308 at 2.0 Å, Asp310 at 3.2 Å, and O1 of xylene at 2.8 Å and O2 at 2.7 Å. The coordination is highly similar to that observed in the A. missouriensis XI (9XIM) mutant E189Q ternary complex with two Mn$^{2+}$ atoms and a linear xylene. It is interesting that in the PirXI structure with the highest activity, which is the Mn$^{2+}$-dependent activity (Table 2), only the M1 and M2b sites are occupied, not M2a.

Xylene was soaked in a Co$^{2+}$-cocrystallized crystal and cryoprotected with 2 M proline. The structure that showed M2–Co$^{2+}$ ion occupies two positions (M2a and M2b) with a 1.8 Å distance (Figure 8B). Furthermore, similar conformational changes for Asp308, Asp310, and Phe61′ are observed as in the Mg$^{2+}$-xylene structure. In this structure, we modeled α-xylene as it was present at a concentration of ~0.25 M in the soaking solution. The position is similar to that of the xylene found in the crystal structure of St. rubiginosus (1XII) and in the neutron diffraction structure of St. rubiginosus (3XWH).

**DISCUSSION**

An efficient xylene isomerase that is well expressed in the active form in S. cerevisiae is critically important for the development of xylene-fermenting yeast strains. The Piromyces sp. E2 xylene isomerase has emerged as an attractive enzyme for this purpose. Our recent work indicates that metal incorporation is critical for in vivo activity, as suggested by the effect of mutations influencing manganese homeostasis on XI activity and anaerobic xylene metabolism. Our work corroborates this by exploring the metal dependence of the catalytic performance and by examining crystal structures of the enzyme soaked with different metals, substrates, and substrate analogues.

Studying the metal dependence of the catalytic activity, we observed that xylene isomerization of PirXI can be activated by various bivalent metals, but the level of activity is dependent on the metals. The enzyme was not activated by a trivalent metal such as Al$^{3+}$. Previous structural studies of XI from Arthrobacter showed that binding of the metal at M2 and substrate binding were disturbed when the enzyme was soaked with Al$^{3+}$ and xylene or fructose. Such a dependence of activity on the type of metal that is bound was also observed in previous studies. It seems that the class I and class II xylene isomerases have different metal preferences as they mostly show the best activity with Mg$^{2+}$ and Mn$^{2+}$ metal cofactors, respectively. As a class II xylene isomerase, PirXI indeed also showed the highest activity with Mn$^{2+}$. Here, we explore the metal dependence of the enzyme by analyzing the structural differences in PirXI reconstituted with different ligands (metals/substrate or analogues) and interpreting the effects on the enzyme activity in view of the proposed catalytic mechanism of xylene isomerases.

The isomerase mechanism includes at least three chemical steps, accompanied and enabled by proton transfer reactions: ring opening of the substrate, hydride transfer from C2 to C1 accompanied by proton transfer from O2 to O1, and ring closure (Figure 1). Structural analysis of xylene isomerase from St. rubiginosus showed that the reaction involves movement of one of the metals (M2, the catalytic metal) and rotation of some of the active site residues such as Asp255 and Lys289, followed by a proton transfer from O2 to O1, and ring closure.
which correspond to Asp308 and Lys342, respectively, in PirXI.26

The His102-Asp105 pair, which is conserved throughout xylose isomerase sequences, is involved in ring opening of the substrate that is bound to M1.26 This ring-opening step seems to be dependent on the metals that are bound to the enzyme. In the structure of PirXI with Cd²⁺ and xylose, the substrate is found in both cyclic and linear form (Figure 6A,B). This indicates a slow ring opening that led to the very low enzyme activity in the presence of Cd²⁺. For many xylose isomerases, Cd²⁺ is reported to inhibit the enzyme reaction, probably by preventing the ring opening of the substrates as the structure with this metal shows only the cyclic form of the substrate.26

After linearization of the substrate M2a that is coordinated with three aspartates, a histidine, and a water molecule shifts to M2b where it now coordinates O1 and O2 of the sugar and must therefore be in the catalytically competent position.26 The mobility of the M2 metal is supposed to aid the reaction after ring opening to facilitate isomerization in its linear form.26,31 A structure of the catalytically most competent enzyme (Mn²⁺ and xylose bound) revealed that Mn²⁺ occupies only the M2b position of the two conformations at the M2 site (Figure 8A). This might indicate that the rapid or more complete shift of the metal from M2a to M2b contributed to the higher activity of the enzyme in the presence of Mn²⁺ compared to the other metals. The kcat of PirXI–Mg²⁺ is half that of PirXI–Mn²⁺, and in the enzyme structure with Mg²⁺, the metal is observed at positions M2a and M2b. In the structure of PirXI–Ca²⁺, the M2 metal is found at only the M2a position (Figure 8A). The large size of Ca²⁺ could hamper its movement, which may be one of the causes of the low rate of reaction of PirXI with this metal.

Variations in transition state stabilization of the actual hydride transfer steps among the different metals bound in the M2b site may also contribute to the observed differences in activity. Movement of the metal at position M2 does not necessarily mean that the metal can activate the enzyme. Although Ni²⁺ was found at both M2 positions, the enzyme showed no detectable activity (Figure S3B). The presence of Ni²⁺ at two distinctive M2 positions was also observed in the structure of PirXI with Cd²⁺, and in the structure of PirXI with Ca²⁺, the metal is observed at M2a and M2b. In the structure of PirXI–Ca²⁺, the M2 metal is found at only the M2a position (Figure 8A). The size of Ca²⁺ could hamper its movement, which may be one of the causes of the low rate of reaction of PirXI with this metal.

PirXI is also active with Yb³⁺ as the metal cofactor, and the specific activity was ~70% of that of PirXI–Mg²⁺. PirXI–Fe²⁺ shows two iron ions in the active site (Figure S3A). Although the reaction of PirXI in the presence this metal was performed under O₂-limited conditions, we cannot exclude the possibility of Fe²⁺ partially being oxidized to Fe³⁺, which would influence the activity. Strong inhibition of a class II XI by iron has been observed in a previous study of another class II XI from Arthrobacter strain B3728, in which 10% of the Ni²⁺ was being oxidized to Fe³⁺, which would in turn influence the enzyme activity. Nevertheless, the property of PirXI makes it more favorable as a specialized catalyst for lignocellulosic bioethanol production as the enzyme will be devoted to only converting xylose to xylulose.

As we have revealed the impact of various metals on the kinetic properties of PirXI, it is important to understand the in vivo metal binding of the enzyme and how it influences its activity. In this study, we have measured the metal content of PirXI isolated from E. coli. The enzyme was found to bind a mixture of several metals, and the contents were different depending on the cultures from which the enzyme was purified. This indicates that the metal contents of the overexpressed enzyme can be influenced by the intracellular metal composition as well as the enzyme–metal affinity. Therefore, measurements of xylose isomerase activities with cell-free extracts or with enzyme samples of which the metal content is not strictly controlled by reconstitution of chelator-treated apoenzyme may give results that have little significance because of variations in metal content, some of which may be introduced during sample preparation. Thus, the medium composition, expression level, and growth phase of cultures may influence metal availability and apparent XI activity.67,69

Previously, Hlima and colleagues found that the recombinant xylose isomerase from Streptomyces sp. SK expressed in E. coli has activity lower than that of the native enzyme, which was explained by the lack of posttranslational modification.71 However, overexpression of the recombinant enzyme could have led to the different metal contents and, hence, the different activities. In a recent study, we have shown that intracellular metal contents play a critical role for the in vivo activity of PirXI expressed in S. cerevisiae as it influences metal loading of the enzyme and, thereby, the in vivo catalytic activity and the rate of xylose metabolism.53
Although the total metal contents of the purified enzyme were close to the theoretical value (two metal ions per monomer of PirXI), it seems that the enzyme loses some of the metals during sample preparation for crystallization. The crystal structure of native PirXI showed that only M1 was occupied, which indicates stronger metal binding at M1 than at M2. Interestingly, His272, one of the M2 metal-coordinating residues, was shown to be doubly protonated half of the time, and this could cause weak binding of the metal, although this could be dependent on the type of metal. The refinement of the structure with different metals at M1 showed that the occupancy of metals decreases in the following order: Ca2+ > Mg2+ > Fe3+. Because the metal content analysis showed that the enzyme bound mainly Mg2+ and lower levels of Ca2+ and Fe3+, we speculate that more Mg2+ was bound at M2 than at M1 in native PirXI. Different affinities of various metal ions for the M1 and M2 binding sites were also observed in previous studies, and the metal preferences differ between the isomerases. For instance, XI from Arthrobacter B3728 showed a high affinity for Mn2+, Co2+, Cd2+, and Pb2+ at M2 and Mg2+ and Al3+ at M1, while XI from St. rubiginosus prefers Mg2+ at position M1 and other various metal ions at position M2.26,61,70

In the study presented here, we measured the activation constants (K_{act}) of different metals, which are probably determined by the binding site with the lower affinity. PirXI has the highest affinity for Mn2+, which is also the superior metal for enzyme activity. The affinity of the enzyme for different metals does not necessarily correlate with the chemical reactivity of the metals for the isomerization reaction. Even though the K_{act} for Co2+ is much lower than that for Mg2+, the enzyme is more active with Mg2+. Interestingly, the metal binding affinity of XI from an Arthrobacter strain is similar to that of PirXI (Mn > Co > Mg), but this enzyme shows the highest activity in the presence of Mg2+. As metal binding residues in xylose isomerases are highly conserved, it is likely that the different chemical reactivity of a metal in different XIs is influenced by the overall protein environment, including second-shell residues and perhaps the residues that are farther from the active site.

The results reported here reveal clear differences between different metals with respect to the binding mode, affinity, and catalytic activity of PirXI. The various assays and crystallography experiments were performed with the apoenzyme reconstituted with different metals. Several factors, including metal-dependent PirXI activity, metal availability, and metal affinity, could affect the efficiency of the enzyme in vivo. It is possible that further exploration using mixtures of metals will be necessary to fully understand the in vivo activity of the enzyme. Previously, the activities of PirXI in the presence of metal mixtures that represented the in vivo metal composition of different S. cerevisiae strains showed that Mn2+ influenced the overall activity significantly, despite it being a minor component of the overall mixtures.33 This suggests a higher affinity of the enzyme for this metal, in agreement with the current half-saturation constants (K_{act}). On the other hand, an inhibitory effect of Ca2+ was not pronounced at concentrations mimicking the in vivo levels, as the activities with the metal mixtures were as high as the activity with Mg2+ alone.

Besides metal content, the pH dependence of XI may be important for the in vivo activity. PirXI shows an optimum pH of ≥7.5 in the presence of Mg2+, Mn2+, and Ca2+. The enzyme activity decreases rapidly at pH <6.5, and this decrease is thought to be related to the local pK_a of His272, which is involved in metal (M2) coordination, or Lys342, which in turn interacts with the metal binding residue Asp310.26 The pH—activity profile seems also to be dependent on the metals as PirXI—Co2+ showed a pH profile different from that of the enzyme with other metals. The activity of PirXI—Co2+ is unaffected by the pH, at least in the range of 5.5—7.5. The pH profiles of xylose isomerases of different biological origin may vary. Some show the best activity in the acidic pH range,26,71—74 and some are most active in the neutral pH range3,69,75 or the basic pH range32,76. Comparing the sequences and structures of these XIs with different pH optima will be useful for engineering PirXI to optimize its activity for the cytosolic pH of the S. cerevisiae.

### CONCLUSIONS

Xylose isomerase from Pirromyes sp. E2, which catalyzes xylose isomerization in metabolically engineered S. cerevisiae strains for conversion of pentoses to ethanol, can accept various metals. The catalytic performance of the enzyme is dependent on the type of metal that is bound, with Mn2+ being the best metal, followed by Mg2+, Ca2+ and Cd2+ poorly activate the reactions, and Ni2+ and Al3+ are not activating. Because there is no clear correlation between the metal-dependent activities and the affinity of the enzyme for metal ions, in vivo XI performance will be influenced by the availability of metal in the medium, uptake and distribution of metal in cells, and incorporation of metal into the enzyme.

Overall, the PirXI crystal structures are similar to other XI structures, the major differences being the N-terminal α-helix extension and the size of the interface area in the tetramers as compared to class I XIs. The active site of each PirXI subunit contains a substrate binding site and two metal binding sites formed by several aspartates and glutamates that are conserved throughout XIs. The results of soaking experiments confirmed that PirXI is highly promiscuous with respect to metal binding and suggest that differences in movement of the catalytic metal to a position required for hydride-shift catalysis and associated conformational changes of metal binding side chains may be related to differences in catalytic performance.

Understanding the metal-dependent catalytic properties of the enzyme through structural and biochemical studies as described in this study provides critical insights for enzyme engineering to optimize the in vivo enzyme activity, thereby improving the production of ethanol from xylose.

### ASSOCIATED CONTENT

#### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.biochem.7b00777.

Sequence alignment of xylose isomerases (Figure S1), active site structures with electron density maps (Figure S2), active site structures of Fe—XI and Ni—XLS—XI (Figure S3), and metal—ligand distances (Table S1) (PDF)

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