The Three-dimensional Structure of Invertase (β-Fructosidase) from Thermotoga maritima Reveals a Bimodular Arrangement and an Evolutionary Relationship between Retaining and Inverting Glycosidases*

Received for publication, December 19, 2003, and in revised form, February 5, 2004
Published, JBC Papers in Press, February 18, 2004, DOI 10.1074/jbc.M313911200

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Invertase, the β-D-fructofuranosidase (EC 3.2.1.26) that cleaves sucrose into fructose and glucose, is one of the earliest discovered enzymes. It was isolated in the second half of the 19th century, and its name was coined because the enzyme produces “invert” sugar, which is a 1:1 mixture of dextrose and levorotatory d-fructose (1). Because of its chemical structure, sucrose can be cleaved by either α-glucosidase or β-fructofuranosidase activity. Koehl and Stein established that invertase is a β-fructofuranosidase by performing the reaction in 18O-labeled water and determining the 18O content of the products (2). The transfructosylation activity of invertase indicated that the enzyme operates with a molecular mechanism leading to overall retention of the anomeric configuration (2). The breakdown of sucrose is widely used as a carbon or energy source by bacteria, fungi, and plants. In plants, both glucose and fructose are implicated in the signaling pathways by which sucrose concentration functions as a key sensor of the nutritional status of plants, and, thus, invertase plays a fundamental role in controlling cell differentiation and development (3, 4). Commercially, invertase is mainly used in the confectionery industry, where fructose is preferred over sucrose because of a sweeter taste and a lower propensity to crystallize.

Although animals, including man, display a strong preference for sucrose-containing diets, their genomes do not encode invertases. Instead, they use a different and unrelated enzyme, sucrose α-glucosidase (EC 3.2.1.48), to hydrolyze sucrose. The genomes of human gut microorganisms such as Bacteroides thetaiotamicron (5) and Bifidobacterium longum (6) do possess invertase genes, demonstrating that these organisms benefit from the large intake of sucrose by humans.

Invertases are found in family GH32 of the sequence-based classification of glycosidases (afmb.cnrs-mrs.fr/CAZY) (7). This family, which includes over 370 members (as of January 2004) from plant, fungal, and bacterial origin, contains not only invertases but also other fructofuranosidases such as inulinase (EC 3.2.1.7), levanase (EC 3.2.1.65), and exo-inulinate (EC 3.2.1.80), and transfructosidases such as sucrose 1-fructosyltransferase (EC 2.4.1.99) and fructan:fructan 1-fructosyltransferase (EC 2.4.1.100).

Glycoside hydrolases or glycosidases are a widespread group of enzymes displaying a great variety of protein folds and substrate specificities. They share a common defining feature in two critically located acidic residues, which make up the catalytic machinery responsible for the cleavage of glycosidic bonds. These two invariant residues have been identified experimentally in yeast invertase as an aspartate located close to the N terminus acting as the nucleophile (8) and a glutamate acting as the general acid/base (9). The enzymatic hydrolysis of glycosidic bonds has two possible stereochemical outcomes, inversion or retention of the anomeric configuration. Invertase is a retaining enzyme (2). With no known exception to date, the molecular mechanism appears conserved among the members of a given sequence-based family (10, 11). Sensitive sequence analyses coupled to structural comparisons have revealed significant similarities between representatives of different families, accompanied by a conservation of the catalytic machinery and of the stereochemical outcome of the reaction, reflecting...
ancient divergence from a common ancestor to acquire novel substrate specificities (12). The evolutionarily, structurally, and mechanistically related families were grouped together in higher hierarchical level termed “clans” (10).

Threading analyses and homology modeling have led to the prediction that, as a member of glycosidase family GH32, invertase would display a six-bladed β-propeller fold related to that of influenza virus neuraminidase (13). However, the recent report on the three-dimensional structure of the family GH68 levansucrase from *Bacillus subtilis* (14) revealed that it had a novel five-bladed propeller fold, which has only been described previously for tachylectin (15) and for the family GH43 α-1-arabinofuranosidase from *Cellvibrio japonicus* (16). Recent detailed sequence analyses have revealed the existence of sequence motifs conserved in the glycosidase families GH32, GH43, GH62, and GH68, suggesting a possible structural relationship between these families (17) despite the opposite mechanisms in GH32 and GH68 (retaining) and GH43 (inverting). It should be noted that, because of the rapid mutarotation of furanosides, it is very difficult to experimentally determine the stereochemical course of the reaction catalyzed by furanosidases such as the family GH43 α-1-arabinofuranosidases. Three independent reports have, however, concluded that family GH43 enzymes operate by an inverting mechanism (18–20).

The mechanism prevailing in family GH62 is not known. After over a century of investigations and almost 40 years since the first crystal structure of a protein was solved, no three-dimensional structure of an invertase or of any member of glycosidase family GH32 has been reported. Here we report the three-dimensional crystal structure of *Thermotoga maritima* invertase. This thermostable enzyme has recently been biochemically characterized by Liebl *et al.* (21), who have determined that it liberates fructose from various substrates such as sucrose, raffinose, and inulin. The structure not only provides a template for all members of family GH32 (including invertases, inulinas, levanas, exo-inulinas, sucros:ar:au:ce 1-fructosyltransferases, and fructan:fructan 1-fructosyltransferases), but it also allows dissection of the exquisite details that distinguish retaining and inverting furanosidases with a perfectly superimposable catalytic machinery.

**EXPERIMENTAL PROCEDURES**

**Protein Cloning, Expression, and Purification—Genomic DNA of *T. maritima* strain MSB8 (DSM 3109), kindly provided by Dr. Wolfgang Liebl (Georg-August-Universität, Göttingen, Germany), was used to amplify the invertase gene (GenBank™ accession number AAD36485).** The *Escherichia coli* strains used were DH5α for cloning experiments and BL21pLysS for expression. Vector pDONR is from Invitrogen, whereas vector pDEST17/O/I is the modified vector pDEST17 from Invitrogen by insertion of lacO and lacI, to prevent expression leakage. The invertase gene was amplified using INV-F (5'-TCTAGACGCAGAATGGATCCTACGTA-3') and INV-R (5'-TCTACACCAATATGTCTCCTCGA-3') primers containing recombination sequences for integration in Gateway™ vectors. PCR was performed using 500 ng of total genomic DNA of *T. maritima*, 300 ng each primer, 1.2 μl of PCR buffer, 1 μl of Pfu polymerase (Invitrogen), 1× Ph buffer (Invitrogen), and 1 mM MgSO4. The amplification program was 94 °C for 5 min followed by 30 cycles of 94 °C for 45 s, 55 °C for 30 s, and 68 °C for 2 min. The amplification was completed with a final extension at 68 °C for 10 min. The amplification product was purified by precipitation in 30% polyethylene glycol 8000 and 3 M LiCl, and re-suspended in 10 μl of TE buffer (10 mM Tris and 0.5 mM EDTA, pH 7.5). The PCR product was cloned in pDONR (Invitrogen) and then in pDEST17/O/I vectors as described in the manual supplied by Invitrogen (22) to obtain the plasmid pINV.

Another single colony of BL21 pLysS containing the pINV plasmid was used to inoculate 40 ml of TBAC broth supplemented with 100 μg/ml ampicillin and 34 μg/ml chloramphenicol. The culture was incubated overnight at 37 °C with constant shaking. The culture was used to inoculate 3 liters of TBAC. Incubation was done at 37 °C with vigorous shaking (240 rpm), and 0.5 μl of isopropyl-1-thio-β-D-galactopyranoside was added when A600 reached 0.8. This induction was followed by another incubation at 37 °C for 4 h. Cultures were pelleted and then suspended in 50 ml of lysis buffer (50 mM Tris pH 8, 150 mM NaCl, 10 mM imidazole, 1 mM EDTA, and 0.1% Triton X-100) containing 1 mM phenylmethylsulfonyl fluoride and 0.25 mg/ml lysozyme. This cell suspension was kept overnight at ~80 °C. After thawing, the lysate was supplemented with 10 μg/ml DNase I and 20 μg/ml MgSO4 and then incubated at 37 °C until it became fluid. The supernatant containing soluble proteins was separated from the pellet by centrifugation (20,000 × g) for 30 min at 4 °C. The SeMet protein was produced as follows. A single colony of BL21 pLysS containing the pINV plasmid was used to inoculate TBAC followed by overnight incubation at 37 °C. This culture was washed several times to remove the traces of TBAC medium and then used to inoculate 1 liter of M9 medium medium from Difco supplemented with 2 mM MgSO4, 0.36% glucose, 100 μM CaCl2, 100 μM ampicillin, and 34 μg/ml chloramphenicol. Incubation was performed at 37 °C under vigorous shaking (240 rpm). When the A660 reached 0.5, 1.5 mM L-lysine, 1.5 mM L-phenylalanine, 1.5 mM L-threonine, 0.8 mM L-leucine, 0.8 mM L-isoleucine, 0.8 mM L-valine, and 0.5 mM seleno-L-methionine (final concentrations) were added. After 30 min of incubation at 37 °C, 0.5 μM isopropyl-1-thio-β-D-galactopyranoside was added to the culture. After induction, expression was followed by measuring A660 until a value of 1.7 was reached. Culture lysis was done as described above.

In all cases, the supernatant of the 20,000 × g centrifugation was filtered (Amicon, 0.2-μm pore-sized membrane), and the invertase was then purified in two steps. First, nickel affinity chromatography was performed using buffers containing 50 mM Tris, pH 8, 200 mM NaCl, and 50 and 500 mM imidazole for the washing and elution steps, respectively. Subsequently, the protein was submitted to gel filtration on a Sephadex column (Amersham Biosciences). The fractions containing the protein were pooled and concentrated to 11 mg/ml for the native thermostable and 8 mg/ml for the SeMet protein over ultrafiltration styrene acrylonitrile membrane (Millipore; cut-off was 30 kDa).

To verify that the N-terminal His-tag did not influence the enzymatic activity of invertase, the hydrolysis of sucrose by the purified protein was monitored. The method employed was adapted from Kidby and Davidson (23) and consisted of the measurement of reducing sugars by ferricyanide. Invertase (200 μl) was incubated at 75 °C in 100 mM sodium acetate buffer (pH 5.5) and 120 mM sucrose, i.e., exactly the same conditions as those described by Liebl *et al.* (21). One hundred-microliter samples were taken at different times of incubation. The enzymatic reaction was revealed by mixing samples with 1 ml of reagent (1 mM K3Fe(CN)6, 130 mM Ca2O4, and 5 mM NaOH) and by heating the samples for 7 min at 85 °C. The activity was monitored by the decrease of A420 as a function of time and led to values (data not shown) very similar to those published by Liebl *et al.* (21).

**Cryocystalization of Native and SeMet-substituted Proteins—Cryostallization conditions were first investigated using two sparse matrix sampling kits (Molecular Dimensions and Stora Forest).** Optimized conditions for obtaining a suitable size were obtained by mixing 15% polyethylene glycol 4000, 150 mM LiSO4, and 100 mM sodium citrate at pH 4.2 with 11 mg/ml native protein. Crystals grew within 3 days at 20 °C by the vapor diffusion method. The conditions for the SeMet-substituted protein were 17% polyethylene glycol 1000, 50 mM LiSO4, 1% isopropanol, and 100 mM sodium citrate buffer at pH 4.2. Here the drops were composed of 2 μl of protein at a concentration of 3 mg/ml with 1 μl of reservoir solution. Both the crystals of native and SeMet-substituted protein belonged to space group P21, with unit cell parameters a = 94.2 Å, b = 113.2 Å, c = 129.6 Å, α = 98.96°, γ = 98.96°. The asymmetric unit contains six monomers giving a Vm value of 2.2 Å3 per Da and 45% solvent content.

**Data Collection and Phase Determination** Crystals were soaked in mother liquor supplemented by 15% glycerol (v/v) before flash freezing in a cryogenic nitrogen stream at 100 K. Diffraction data of native and SeMet-substituted protein crystals, both in space group P21, were collected at the European Synchrotron Radiation Facility (ESRF, Grenoble, France) on beam lines ID14-E2H and ID29 respectively (Table 1). The data on native and SeMet-substituted crystals were processed in the abovementioned peak (space group P21) and phased using the SAD method. Forty of the 38 SeMet positions were determined by anomalous Patterson maps using the subroutine XPREP of the program package SHELEX5,0 (24). The 40 sites were refined with SHARP (25), and the missing eight positions were found in the residual maps. The 48 selenium positions appeared to be arranged in a manner that suggested the presence of three dimers. The asymmetric unit contains six monomers giving a Vm value of 2.2 Å3 per Da and 45% solvent content.
The crystal structure of the

**RESULTS AND DISCUSSION**

**Overall Fold**—The crystal structure of the *T. maritima* invertase (residues 1–432) has been solved by SAD phasing of the SeMet-substituted protein at a maximal resolution of 2 Å. The SeMet-substituted as well as the native crystals belong to space group P2₁, with unit cell parameters a = 94.2 Å, b = 113.2 Å, c = 129.6 Å, and β = 98.96°. The coordinates describing six copies of the invertase polypeptide chain and 1754 water molecules per asymmetric unit were refined to final R- and Rfree-factors of 17.6 and 22%, respectively. One molecule of invertase is composed of two individual modules, namely a five-bladed β-propeller (residues 1–295) catalytic module linked to a C-terminal β-sandwich module (residues 306 to 432) by a 10-residue linker (Fig. 1). The ensemble of six bi-modular molecules arrange into three individual dimers, displaying 2-fold symmetry each. The three dimers are not related by any point group symmetry but by non-symmetrical rotations and translations. The dimer arranges around a pseudo 2-fold axis, bringing the β-sandwich domain of monomer A in contact with the β-propeller domain of one of the six invertase molecules. Ramachandran statistics (PROCHECK) indicated that, for the overall structure of the six molecules present in the asymmetric unit, 87.1% of the atoms are in the most favored region, and 12.6% are in additionally allowed regions. Details of refinement statistics are summarized in Table I.

![Summary of data collection, phasing, and refinement statistics](image)

| Data sets | Native | λ <sub>1</sub> (Peak) |
|-----------|--------|----------------------|
| Wavelength (Å) | 0.979 | 0.97904 |
| High resolution (Å) | 2.0 | 2.2 |
| (Anomalous) completeness (%) | 99.9 (99.9) | 99.3 (98.6) |
| Redundancy | 3.1 (3.1) | 4.2 (4.2) |
| I/σ(I) | 8.0 (2.1) | 6.8 (2.2) |
| R<sub>sym</sub> | 0.077 (0.402) | 0.073 (0.211) |
| Phasing statistics | | |
| Anomalous difference (%) | | |
| Figure of merit (overall) | 0.429 (0.853) | 6.4 |

* Numbers in parentheses indicate values for the highest resolution bin.

<sup>a</sup> R<sub>sym</sub> = ∑<i>[I</i> –<i>I</i>] / <i>I</i>, where <i>i</i> is the ith measurement and <i>I</i> is the weighted mean of <i>I</i>.<br>
<sup>b</sup> Figure of merit value in parentheses is calculated after density modification with the DM program.<br>
<sup>c</sup> R<sub>cryst</sub> = ∑<i>[F</i><sub>obs</sub> –<i>F</i>]<i>/F</i><sub>obs</sub>.<br>
<sup>d</sup> R<sub>free</sub> is the same as R<sub>cryst</sub> for 5% of the data omitted from refinement totaling 10,694 reflections.<br>
<sup>e</sup> R.m.s. is root mean square.
FIG. 2. Sequence alignment of a selection of family GH32 invertases. The sequences are identified as follows: Tmar_inv, *T. maritima* invertase (Swiss-Prot O33833); Ecol_inv, *E. coli* K12 invertase (Swiss-Prot P16553); Smut_inv, *Streptococcus mutans* GS-5 invertase (Swiss-Prot P13522); Zmai_inv, *Zea mays* invertase (Swiss-Prot O81189); Atha_inv, *Arabidopsis thaliana* Landsberg erecta (GenBankTM BAA89048.1); Scer_inv1, *Saccharomyces cerevisiae* invertase 1 (Swiss-Prot P10594); and Scer_inv4, *S. cerevisiae* invertase 4 (Swiss-Prot P10596). The boxes shaded in red are strictly conserved residues, whereas the boxes shaded in light blue concern highly similar sequence regions. The sequence numbering and secondary structure elements (the color codes of the secondary structure elements are the same as in Fig. 1) correspond to the sequence of *T. maritima* invertase. The highly conserved motifs A through F, as defined by Pons et al. (13), are highlighted by left and right arrows above the sequences. The alignment was produced with ClustalX (46), and the figure was produced with ALSCRIPT (47).
The catalytic site at the center of the sugar binding site of T. maritima invertase has an elliptical shape with approximate dimensions of 63 × 43 × 45 Å with a negatively charged surface depression at the center of the β-propeller.

The clearly defined electron density revealed two amino acid residues in conflict with the GenBank™ sequence (A108 → V108 and V179 → A179). Therefore, the nucleotide sequence was checked twice (amplification from genomic DNA and the expression clone), and the two single base differences (C323 → T332 and T536 → C536) were only detected for the expression clone. As a consequence, these mismatches are attributed to misincorporation by the polymerase Pfx. Nevertheless, activity tests (see “Experimental Procedures”) indicated that these mutations do not affect the enzymatic activity.

A five-bladed β-propeller structure has first been reported for tachylectin (15) and was found more recently for the enzymes α-1-arabinanase (16) and levansucrase (14) of the glycoside hydrolase families GH43 and GH68, respectively. Highly similar to the families GH43 and GH68 structures, the five β-sheets of invertase, labeled I–V (Fig. 1), adopt the classical “W” topology of four antiparallel β-strands. The N-terminal second strand lines the central cavity, and the C-terminal last strand is at the periphery, to which the β-sandwich module is connected by a short linker. Interestingly, and in contrast to levansucrase and α-1-arabinanase, the five bladed β-propeller of invertase does possess the short “molecular Velcro” that is typical of six- and seven-bladed β-propellers (15, 34, 35). The N-terminal first strand forms the outermost β-strand of the C-terminal blade V; however, only one hydrogen bond is formed across the sheet (Phe-8 O-Met-277 N, 2.8 Å). A similar short Velcro has also been observed in the six bladed β-propeller of Vibrio cholerae sialidase (36). As in all β-propeller structures, the β-strands forming the blades are strongly twisted, giving an angle of ~90° between the first and last β-strand of a blade. Insertions are common in this type of β-propellers, and, likewise, short stretches of 3-10-helices are found inserted between several individual β-strands of the structure described here. They are, however, less extended than in the GH68 levansucrase, and from this perspective the β-propeller of invertase resembles more that of GH43 α-1-arabinanase.

The Catalytic Active Site—The catalytic active site is positioned at one end of the cavity at the center of the β-propeller with a funnel-like opening toward the molecular surface. It clearly has a pocket topology, which is fully consistent with the strict exo mode of action of the enzyme on the fructose polymer inulin (21). The three carboxylate groups of two aspartate (Asp-17 and Asp-138) residues and one glutamate (Glu-190) residue point to the center of the depression and generate a high negative charge at the active site. Reddy and Maley have shown that Asp-23 in yeast invertase (Asp-17 in T. maritima invertase) is the catalytic nucleophile (8), whereas Glu-204 (here Glu-190) is the general acid/base (9). In addition to the two regions containing the catalytic machinery, multiple sequence alignments of the GH32 family (Fig. 2) have revealed a number of other highly conserved amino acid stretches (13, 37). The inspection of the three-dimensional structure allows us to define possible roles for these highly conserved residues. For the family GH68 levansucrase, the sucrose complex of an inactive mutant has also been reported (14). Because the catalytic modules of invertase and levansucrase are structurally related, the superimposition of invertase with the sucrose-containing complex of levansucrase (PDB identification code 1PT2) allows us, by similarity, to infer the position of a sucrose molecule and model it in the active site of invertase (Fig. 3A). The crystal structure of invertase revealed a glycerol molecule, present in the substrate binding site, that mimicked the O4’ and O6’ hydroxyl groups of the substrate fructose moiety (Fig. 3B), and this helped define the precise position of the modeled sucrose molecule. This model shows that the second strictly conserved aspartate residue in motif D (for motif definitions see Ref. 13

| Table II | Hydrogen bonding and close contacts between modeled sucrose and invertase active site residues |
|----------|------------------------------------------------------------------------------------------|
| Sucrose atom | Invertase residue | Distance |
| Fructose O1’ | Asp-17-O’1 | 2.9 |
| Fructose O2’ | Asp-17-O’2 | 3.4 |
| Glucose O4 | Asp-17-O’2 | 3.4 |
| Asp-236-N’2 | 3.5 |
| Fructose O3’ | Asp-17-O’3 | 2.9 |
| Glucose O2’ | Glu-190-O’2 | 3.6 |
| Asp-138 | 2.9 |
| Asp-138-O’3 | 3.3 |
| Fructose O4’ | Asp-138-O’4 | 2.5 |
| Fructose C6’ | Asp-138-O’4 | 2.6 |
| Fructose O6’ | Ser-75-N’1 | 3.0 |
| Glucose O1 | Arg-137-N’1 | 3.2 |
| Glucose O2 | Arg-137-N’1 | 3.8 |
| Glucose O4 | Arg-137-N’1 | 3.8 |
Asp-138 in *T. maritima* invertase, forms hydrogen bonds to O3 and O4 of the fructose unit, whereas the neighboring Arg-139 is hydrogen-bonded to the glucose O4. Apparently, the pair of strictly conserved residues, "RD," binds to characteristic hydroxyl groups of the substrate and, therefore, most likely plays a crucial role in substrate binding and recognition. Interestingly, the enzymes of family GH68, which hydrolyze the same substrates, also have the highly conserved motif "RDP," whereas GH43 and GH62, which have a structurally related fold but hydrolyze different substrates, do not possess this motif and only have the aspartate residue in the same position. The highly conserved motif designated A by Pons *et al.* (13) contains the nucleophile Asp-17 and the preceding Asn-16, which forms a hydrogen bond to the O6 group of fructose, whereas the sequence regions designed B and B1 appear to be structurally important, because the conserved aromatic residues are involved in hydrophobic interactions in the face-to-face packing of blades I and V and are not in the catalytic site. However the side chain of Trp-41, located between motifs B and B1, points into the active site and is most probably part of the aglycone binding pocket. Motif C contains residues involved in substrate binding such as Phe-74, which borders the fructose binding pocket, and Ser-75, which forms hydrogen bonds to the O4 hydroxyl of fructose (3.5 Å) and to catalytic nucleophile (2.9 Å). The sequence region E contains the general acid/base Glu-190 (3.1 Å from the glycosidic oxygen) and Cys-191, both located in the heart of the active site. This conserved cysteine is most probably important for transition state stabilization and/or the catalytic residue microenvironment, because it forms hydrogen bonds to Asp-17 (3.5 Å) and Asp-138 (3.6 Å).
is interesting to note that enzymes of family GH-68 have an arginine replacing this cysteine, although they cleave highly similar substrates. The importance of these differences for binding, catalysis, and stability will be investigated in the future by a study of inactivated invertase mutants in complex with oligosaccharides. See Table II for a comparison of hydrogen bonding and close contacts between modeled sucrose and invertase active site residues.

**Structural Relationship to Families GH68 and GH43 Five-bladed β-Propellers**—Based on detailed sequence analyses, a structural relationship between families GH32, GH43, GH62, and GH68 has been predicted (13, 17). The common five-bladed β-propeller fold, recently revealed by the structural determinations of members of family GH68 (14) and GH43 (16), confirmed this structural relationship. The crystal structure of invertase now proves that the catalytic modules of family GH32 enzymes also display the same five-bladed β-propeller fold. The superimposition of the catalytic module of invertase onto the two other enzymes leads to an overall root mean square deviation of 3.24 Å for 306 Cα atoms in the case of the family GH32 α-L-arabinanase and 3 Å for 359 Cα atoms in the case of the family GH68 levansucrase. Whereas levansucrase and invertase both retain the anomic configuration at the site of cleavage, α-L-arabinanase is an inverting enzyme (18–20). The most widely accepted (and documented) view of the difference between the catalytic machineries of retaining and inverting glycosidases is that, in the former, the two catalytic amino acids are -5.5 Å apart, and in the latter this distance is generally -9 Å, with the exception of β-helical enzymes such as polygalacturonase or α-carrageenase (38, 39). Remarkably, the three invariant amino acids Asp-17, Asp-138, and Glu-190 in GH32, defined as the catalytic residues in each of the families GH32, GH68, and GH43, superimpose rather well in all three enzyme structures (Fig. 4A), showing that the relatedness is not solely with the fold but also with the catalytic machinery. The structural superposition shows that there is no difference in the distances of the catalytic residues relative to each other, as has generally been observed in inverting versus retaining glycoside hydrolases (10, 12, 40, 41). Instead, it is the difference in the binding position of the sugar in the -1 subsite (subsite nomenclature of Davies et al.; Ref. 48) that makes the difference in the catalytic mechanism of invertase and levanase on the one hand and α-L-arabinanase on the other. The arrangement of the loops and residues surrounding the catalytic machinery in α-L-arabinanase is such that the arabinosyl moiety in the -1 subsite is bound in a position almost perpendicular to the fructofuranosyl moiety in invertase and levanase. Consequent to this different binding, the nucleophilic residues are only -3.6 Å from the sugar C1 atom in invertase and levanase (14), whereas the distance C1-Asp-38 in α-L-arabinanase is 6 Å, leaving room for a water molecule (16) (Fig. 4B). This different binding mode of the “glycone” part of the substrate fully explains the opposite stereochemic outcome of the reaction, despite a perfectly superimposable catalytic machinery.

**The β-Sandwich Module**—The C-terminal residues (from 306 to 432) of *T. maritima* invertase compose an individually folded β-sandwich consisting of two sheets of six β-strands. This module is connected to the catalytic module via a short, 10-residue-long linker region that is wrapped around the β-sandwich. Contrary to the catalytic module, which can be readily aligned with all other members of glycosidase family GH32, BLAST searches conducted with the C-terminal module of *T. maritima* invertase did not reveal a statistically significant sequence similarity with the equivalent regions in other family GH32 proteins. To detect possible relatedness beyond the detection level of BLAST, we have removed the easily identifiable catalytic domain region in all complete family GH32 members and constructed a sequence library with the remaining C-terminal regions. PSI-BLAST searches conducted starting with the C-terminal region of plant or fungal or bacterial family GH32 members picked the *T. maritima* C-terminal domain after a few iterations, indicating that all GH32 family members will also be appended to a β-sandwich domain, such as that of *T. maritima* invertase.

The alignment of this module with the programs DALI (42) and 3D-PSSM (43) onto other β-sandwich structures revealed structural similarities with the β-sandwich in galectins, the Charcot-Leyden crystal protein, carbohydrate binding modules (CBMs), and other more distant proteins like lectins and exotoxin A. The highest similarity is observed with the human galectin-3 (Protein Data Bank identification, 1AK3; DALI Z-score, 10.9; root mean square deviation for 127 Cα is 2.4 Å) (Fig. 5) and with the Charcot-Leyden crystal protein (Protein Data Bank identification, 1CLC; DALI Z-score, 10.7; root mean square deviation for 132 Cα is 2.6 Å), which has recently been found to be a maltose binding galectin (44). It is interesting to note that six-bladed β-propeller glycosidases such as *E. coli* aminoglycosidases have also been found appended with lectin-like domains (36, 45). It has been observed that extracellular yeast invertase, a functionally active homodimer in solution, acquires maltose to self-assemble into higher oligomers upon transport and secretion (33). It is therefore tempting to postulate that the supplementary β-sandwich module of yeast invertase plays the role of a carbohydrate recognition domain involved in the higher oligomer formation. The distant similarity of the C-terminal module of *T. maritima* invertase, compared with the other members of the GH32 family, suggests that this module has perhaps lost this function in *T. maritima* invertase. Alternatively, this module might have evolved in *T. maritima* invertase to preserve stability at high temperature, even if the ancestral function of it has been lost. Proteins from hyperthermophilic organisms frequently adopt a modular as well as a multimeric structure. These two complementary features can on July increase stability at high temperature by masking weak regions at the surface of the protein.

**Acknowledgments**—We thank Dr. Wolfgang Liebl (Georg-August-Universität, Göttingen, Germany) for his generous gift of *T. maritima* genomic DNA. We also thank the staff of the European Synchrotron Radiation Facility for the provision of beam time and for technical assistance at the beamlines ID29 and ID-14 EH2. We thank Dr. J. Allouch and Dr. A. Gruez for helpful discussions.

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J. Biol. Chem. 2004, 279:18903-18910.
doi: 10.1074/jbc.M313911200 originally published online February 18, 2004

Access the most updated version of this article at doi: 10.1074/jbc.M313911200

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