Three-dimensional structure of *Schistosoma japonicum* glutathione S-transferase fused with a six-amino acid conserved neutralizing epitope of gp41 from HIV

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Abstract

The 3-dimensional crystal structure of glutathione S-transferase (GST) of *Schistosoma japonicum* (Sj) fused with a conserved neutralizing epitope on gp41 (glycoprotein, 41 kDa) of human immunodeficiency virus type 1 (HIV-1) (Muster T et al., 1993, *J Virol* 67:6642-6647) was determined at 2.5 Å resolution. The structure of the 3-3 isozyme rat GST of the ß gene class (Ji X, Zhang P, Armstrong RN, Gilliland GL, 1992, *Biochemistry* 31:10169-10184) was used as a molecular replacement model. The structure consists of a 4-stranded /β/-sheet and 3 α-helices in domain 1 and 5 α-helices in domain 2. The space group of the Sj GST crystal is P432_12, with unit cell dimensions of a = b = 94.7 Å, and c = 58.1 Å. The crystal has 1 GST monomer per asymmetric unit, and 2 monomers that form an active dimer are related by crystallographic 2-fold symmetry. In the binding site, the ordered structure of reduced glutathione is observed. The gp41 peptide (Glu-Leu-Asp-Lys-Trp-Ala) fused to the C-terminus of Sj GST forms a loop stabilized by symmetry-related GSTs. The Sj GST structure is compared with previously determined GST structures of mammalian gene classes ß, ß, and ß. Conserved amino acid residues among the 4 GSTs that are important for hydrophobic and hydrophilic interactions for dimer association and glutathione binding are discussed.

Keywords: fusion protein; glutathione S-transferase; human immunodeficiency virus type 1; *Schistosoma japonicum*; 3-dimensional structure

Glutathione S-transferases (GSTs) are a family of multifunctional enzymes that catalyze the nucleophilic addition of the reduced thiol of glutathione to a variety of electrophiles (Mannervik & Danielson, 1988; Armstrong, 1991; Gilliland, 1993; Rushmore & Pickett, 1993). They are important enzymes involved in the metabolism of potentially toxic alkylating agents. Cytosolic GSTs, dimeric proteins with a molecular mass of each subunit of about 26 kDa, are also involved in intracellular binding and transport of hydrophobic ligands and participate in the synthesis of prostaglandins and leukotrienes. Specific molecular forms of GST are known to be expressed in the cellular mechanisms of drug resistance (Tsuchida & Sato, 1992).

GSTs from a variety of organisms have been identified and characterized (Hiratsuka et al., 1990; Dominey et al., 1991; Meyer et al., 1991; Mannervik et al., 1992; Trottein et al., 1992; Mignogna et al., 1993), including the detection in a wide range of parasitic helminths (Brophy & Barrett, 1990; Brophy et al., 1990). Cytosolic GSTs of the genus *Schistosoma* participate in the immunogenicity to the vertebrate host and have been suggested as potential components of a vaccine against schistosomiasis (Capron et al., 1987; Sher et al., 1989). In addition, the development of a gene expression system employing the GST from *Schistosoma japonicum* (Sj) allows the fusion of a protein or protein fragment to the C-terminus of the enzyme (Smith & Johnson, 1988; Fainsod et al., 1991; Menéndez-Arias et al., 1992; Oshima-Hirayama et al., 1993; Slany et al., 1993). The fusion peptide or protein is coupled to GST through a linker that can be cleaved by a specific protease, thus providing a useful sys-
tem for cloning, expressing, and purifying the desired protein or peptide product by glutathione-based affinity chromatography.

A number of crystal structures of mammalian GSTs complexed with a variety of ligands in the active site have been recently determined (Reinemer et al., 1991, 1992; Ji et al., 1992; Sinning et al., 1993; Garcia-Sáez et al., 1994). The structures have revealed a common folding pattern with a number of variations in the binding mode of glutathione and especially the xenobiotic substrate binding site. Crystallographic studies have also been instrumental in determining the structural features of the active site essential for catalysis and substrate specificity (Liu et al., 1992, 1993; Manoharan et al., 1992; Ji et al., 1993, 1994).

In this article, the 3-dimensional crystal structure of Sj GST with a C-terminal fusion peptide and reduced glutathione in the active site is described and compared to the 3 previously determined GST structures from the mammalian gene classes, α from human (Sinning et al., 1993), α from rat (Ji et al., 1992), and π from pig (Reinemer et al., 1991). Because Sj GST shares a striking sequence homology with the mammalian α GST (Smith et al., 1986; Hughes, 1993), this structural comparison reveals the shared characteristics of the amino acid residues in the glutathione binding site of both the vertebrate and invertebrate GSTs. The sequence of the fusion peptide is Glu-Leu-Asp-Lys-Trp-Ala from gp41 (glycoprotein, 41 kDa) of human immunodeficiency virus type 1 (HIV-1). Muster and coworkers (1993) determined that this peptide fused with GST was bound to the human monoclonal antibody (Mab) 2F5. This sequence of gp41 was found to be highly conserved in a variety of HIV-1 isolates, suggesting its potential crossreactivity to the antibodies with properties similar to Mab 2F5. The original goal of this work was to crystallize the complex of Sj GST fusion protein with Mab 2F5, but only a crystal of the GST fusion protein was obtained. Nevertheless, because a peptide or protein fragment by itself is often difficult to crystallize, the GST fusion protein represents a great potential for the structure determination of smaller protein fragments. The significance of this fusion method is discussed further elsewhere (Carter et al., 1994).

Results and discussion

Of the 4 GST structures compared, only Sj GST has a monomer in its crystal asymmetric unit. For the others, the 2 monomers in a dimer-paired asymmetric unit are called subunits A and B. Following this practice, the Sj GST monomer in the asymmetric unit is also designated subunit A, and its symmetry-related counterpart subunit B. Unless specified otherwise, all structural comparisons were made with subunit A of each GST. The atomic coordinates of α, π, and γ GSTs were obtained from Brookhaven Protein Data Bank (identification codes 1GST, 1GUH, and 1GSR, respectively). Furthermore, the amino acid residue names and numbers used in this article are for Sj GST; for the others, residue names and numbers are used in conjunction with a specific GST. The coordinates of the Sj GST structure are deposited in the Protein Data Bank (identification code 1GNE).

Final model of the structure

The crystal structure of Sj GST with the gp41 fusion peptide at 2.5 A resolution is shown in Figure 1 and Kinemage 1. It is a monomer forming a perfect dimer by crystallographic 2-fold rotation (Fig. 2). It shares similar characteristics with other GSTs: domain 1 (residues 1-84) is a typical GST folding pattern of βαβαββ, and domain 2 (residues 85-217) consists of 5 α-helices followed by a long stretch of loops (Fig. 3). The summary of least-squares refinement parameters is shown in Table 1. Also, the φ-ψ torsion angle plot for main-chain atoms is shown in Fig-
Schistosoma japonicum glutathione S-transferase fusion protein

Table 1. Summary of least-squares refinement parameters of Sj GST fusion protein complexed with glutathione

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Target value</th>
<th>Final model</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reflections from 6.0 to 2.3 Å with I &gt; 1σ(I)</td>
<td>8,917</td>
<td></td>
</tr>
<tr>
<td>Reflections from 6.0 to 2.5 Å with I &gt; 1σ(I)</td>
<td>7,622</td>
<td></td>
</tr>
<tr>
<td>Crystallographic R* factor (6.0-2.3 Å)</td>
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<td></td>
</tr>
<tr>
<td>Crystallographic R factor (6.0-2.5 Å)</td>
<td>0.219</td>
<td></td>
</tr>
<tr>
<td>RMS deviation from ideal distances (Å)</td>
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<td></td>
</tr>
<tr>
<td>Bond distances</td>
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<tr>
<td>Angle distances</td>
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<td>0.038</td>
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<tr>
<td>Planar 1-4 distances</td>
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<tr>
<td>RMS deviation from bond angle (°)</td>
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<tr>
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</tr>
<tr>
<td>RMS deviation from ideal chirality (Å²)</td>
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<td>0.269</td>
</tr>
<tr>
<td>Thermal parameter correlations (mean/Δβ)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Main-chain bend</td>
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<td>1.300</td>
</tr>
<tr>
<td>Main-chain angle</td>
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<tr>
<td>Side-chain bend</td>
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<td>1.269</td>
</tr>
<tr>
<td>Side-chain angle</td>
<td>1.500</td>
<td>1.600</td>
</tr>
</tbody>
</table>

* R = \( \sum_{\text{all}} |F_o| - |F_c| / \sum_{\text{all}} |F_o|.

Fig. 3. Amino acid sequences of GSTs. The alignments are based on structural considerations. Secondary structural elements are marked by enclosed lines, and common residues among all GSTs are shaded. The Sj GST sequence continues with those of linker peptide and the gp41 fusion peptide.

Fig. 4. The non-glycine residues with near-unfavorable torsion angles are Gln 66, which is involved in glutathione binding, Asn 143, which is in the loop between the α-helices 5 and 6, and Glu 227 in the fusion peptide. There are 2 cis-prolines at residues 55 and 201. Furthermore, a highly ordered glutathione structure was determined in the binding site (Fig. 5). The asymmetric unit has 126 water molecules, 30 of which do not have direct interactions with the protein itself but with other water molecules. Most of the water molecules are found in the open region at the dimer interface, around the glutathione binding site, and around the β-strands in domain 1.

Omit maps showed that the residues of the main Sj GST structure are well defined. These residues have low temperature factors (4-15 Å²) (Fig. 6). Temperature factors for the fusion peptide are relatively higher (15-30 Å²). Electron density for the linking amino acids Leu 220 and Val 221 is not clearly defined; these are the residues with hydrophobic side chains that are in the bulk solvent region. The electron density map indicates the possibility of alternative conformations for these residues.

The gp41 fusion peptide is extended by the linker peptide (Ser-Asp-Leu-Val-Pro-Arg-Gly-Ser-Met; Fig. 3; see Kinemage 1) to
interact with the symmetry-related GSTs (Fig. 7) and consists of a series of tight turns (Fig. 8). The loop formed by the residues 224-229 is stabilized by the interactions between Ser 225 and Asp 229 and a water molecule forming hydrogen bonds with the carbonyl O atoms of Met 226 and Glu 227 and the side chain of Asp 229. In addition, the side chain of Met 226 is positioned between those of Leu 162 and Tyr 163 of the symmetry-related Sj GST. The relative increase in temperature factors for the C-terminal residues and fusion peptides reflects the higher mobility of this portion of the structure (Fig. 6). The same trend was also found in the refined structure of μ GST.

Comparison of amino acid sequences and tertiary structures

Sequence alignment of Sj GST and rat μ GST resulted in 92 residue pair identities, which is 92/217 = 42% (Fig. 3). Also, there are amino acid pairings that may be considered homologous: (1) Leu and Ile in 7 cases, (2) Glu and Asp in 5 cases, and (3) Lys, Arg, and His in 6 cases. Thus, the effective sequence homology is (92 + 18)/217 = 51%. This percentage could be greater with more generous criteria, such as residues with similar polar side chains. The corresponding values for the comparison between Sj GST and α GST are 24% and 33%, and between Sj GST and π GST are 27% and 32%.

The sequence alignment between Sj GST and μ GST shown in Figure 3 also indicates the switching of side-chain polarities: (1) positive and negative side chains in 5 cases, (2) positive and polar residues in 11 cases, and (3) negative and polar residues in 6 cases. These changes are accompanied by compensating interactions. For example, at the interface of the dimer pair, the side chain of Asp 76 from subunit A interacts with the side chain of Arg 88 of subunit B, whereas in μ GST, the amino acid residues are changed to Arg 81 and Asp 97, respectively.

All 4 GSTs share similar structural characteristics: βαβαβα folding pattern in domain 1, and primarily α-helices in domain 2. To compare the tertiary structures, positional and rotational differences between domain 1 and domain 2 were made by su-
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![Graph](image)

**Fig. 6.** Plot of temperature factors of main-chain atoms N, Cζ, and C of Sj GST with the fusion peptide. Bars at the bottom of the graph represent the secondary structural elements: thick bars for β-strands and thin bars for α-helices.

Fig. 7. Stereo view of a Cα tracing of Sj GST with its fusion peptide interacting with symmetry-related GSTs.

The difference in Cα atom distances between the α-helices 4 and 6 is about 1.2 Å between α GST and the other GSTs. The rotation angle is also larger by about 2° when α GST is compared with the other 3 GSTs. Furthermore, another source of the structural difference is that the α-helix 4 of Sj GST is curved by about 15°, whereas that of μ GST is almost linear.

**Hydrophobic and hydrophilic interactions**

A common hydrophobic interface between the dimer pair can be observed in all 4 GSTs. The residue in the loop after the α-helix 2 is the conserved Phe 51 and nearby are the residues Leu 64 and Ala 69 that are at the end of the β-strand 4 (Fig. 3). These residues face the residues in subunit B of the dimer pair, Ala 89, Met 93, Leu 94, and Phe 132. The sequence number and identity of residues that participate in this hydrophobic interaction differ slightly in the other GSTs, but the above-mentioned residues make up the core structure for the hydrophobic interaction between the GST dimer pairs. The region along the dimer 2-fold axis is a relatively open hydrophilic channel, and flanking this channel on both sides of the dimer is this hydrophobic interface acting as a stabilizing force for the GST dimer.

The hydrophilic interactions between the dimer pair is more variable than the hydrophobic interactions. Still, certain key sequence positions can be distinguished: Gln 66, Arg 72, Asp 76, Arg 88, Ser 92, and Asp 100. These residues are located in the α-helices 3 and 4, and together with the corresponding α-helices of subunit B, they are positioned nearly parallel along the 2-fold axis and form a hydrophilic core. The residue Gln 66 participates in the glutathione binding and interacts with Asp 100 of subunit B. The residue Arg 72 interacts with its own symmetry-related residue in subunit B. The residue Asp 76 interacts with Arg 88 of subunit B. In μ GST, the polarity is changed so that Arg 81 interacts with Glu 90 and Asp 97 of subunit B. In π GST, Arg 77 interacts with Asp 88 of subunit B. The residue Ser 92 shows no direct interaction with subunit B in Sj GST; however, in the
other 3 GSTs, this sequence position is aspartate, which shows interactions with a positively charged residue in subunit B.

There are other hydrophilic residues that are conserved in all 4 GSTs at key positions in the sequence. One of these is the residue Asp 151, which is at the beginning portion of the α-helix 6. Its side-chain atoms are observed to interact with the main-chain N atoms of Leu 142 and Thr 148, which are in the loop region after the long α-helix 5. It would seem that the role of Asp 151 is to stabilize the loop region between 2 α-helices. Indeed, a site-directed mutagenesis study showed that the aspartate at this position in ρ GST is important for thermostability of the protein (Kong et al., 1993). Another amino acid with a similar configuration is Glu 87; it is at the beginning portion of α-helix 4, and its side chain interacts with the N atom of Gly 83, which is in the 78–84 loop connecting the 2 domains of GST. Furthermore, aspartate residues at the θ-turn in domain 1 are observed to make interesting interactions with other residues. The side chains of Asp 59 and Lys 77 interact with each other, and Asp 61 is close to Lys 86 of subunit B.

The C-terminal region of Sj GST (residues 213–217) essentially sits on the surface of domain 2. It is also close to the 33–35 loop, a shortened version of the 33–42 loop of μ GST, that links the β-strand 2 and α-helix 2 in domain 1. The amino acid residues of these 2 regions interact closely with each other. In Sj GST, the side chain of Glu 33 forms a hydrogen bond with the ring atom Nε1 of Trp 205; in μ GST, the analogous interaction is between Asp 36 and Lys 210, though in this case Asp 36 is 2 residues farther down both in sequence and spatial relationship than Glu 33 of Sj GST in the interaction with domain 2. The next residue in Sj GST is Arg 34, and its side chain interacts with the side chain of Asp 213 and the carbonyl O atom of Gly 212. The corresponding residue in μ GST is Met 34, and rather than its side chain pointing outward as that of Arg 34 of Sj GST, it faces into the protein core. The 33–42 loop of μ GST interacts with its own symmetry-related loop and residues of domain 2. In π GST, although the corresponding loops have similar conformations, they are farther apart than in Sj GST. This allows the symmetry-related GST to move into this gap. In α GST, the C-terminal α-helix 9 points toward the glutathione binding site and runs nearly parallel with the α-helix 2. There are hydrophobic interactions between Ile 218, Phe 219, Val 54, Leu 40, and Ile 34. The hydrophilic interaction between Arg 220 and Asp 41 is also observed. Unlike μ GST and π GST, α GST does not have a close symmetry-related interaction at this junction; it would seem that the α-helix 9 is sufficient to close the gap between the 2 domains of GST. Thus, either the close interactions between the 2 domains or crystal packing at this region encloses the glutathione binding site.

Glutathione binding site

In all 4 GSTs, an extended form of glutathione is bound to domain 1. Generally, there are 4 types of interaction with glutathione in the binding site (Figs. 5B, 10): (1) stabilization and orientation of the γ-glutamate of glutathione, (2) alignment of the glutathione peptide backbone, (3) stabilization of the terminal carboxylate of glycine, and (4) interaction with the sulfhydryl of cysteine for enzymatic activation. The binding site residues that are common in all 4 GSTs are Tyr 6, Asn 53, Leu 54, Gln 66, and Ser 67 in subunit A, and Asp 100 of subunit B (Fig. 10A; Kinemages 1 and 2).

Association and orientation of the γ-glutamate is accomplished by either a Ser-Gln or Thr-Gln sequence that form hydrogen bonds with the carboxyl group (atoms Oα1 and Oα2). With the exception of π GST, this configuration is further stabilized by a salt bridge between the atom N of γ-glutamate and Asp 100 of subunit B. In π GST, the γ-glutamyl carboxylate group is rotated approximately 180° primarily by a more favorable salt bridge with the side chain of Arg 13. The arginine residue at this sequence location is also present in α GST. In contrast, leucine is the residue in Sj GST and μ GST (Fig. 3).

Hydrogen bonds to the glutathione peptide backbone are produced by another important dipeptide sequence, Leu-Asn in Sj GST and μ GST, Val-Gln in α GST, and Leu-Gln in π GST, and
Fig. 9. Stereo views of Cα atoms of Sj GST (in thick lines) with other GSTs (in thin lines) after the superposition of domain 1.
A: μ GST. B: α GST. C: π GST.

influenced by the presence of Trp or Phe residues. In Sj GST and μ GST, the carbonyl atom Oω of glutathione cysteine forms a hydrogen bond with the indole ring atom Ne1 of Trp 7, and the backbone N atom of glutathione cysteine interacts with the Leu-Asn sequence. In contrast, the presence of Phe in α GST and π GST causes the peptide flip of glutathione cysteine, so that the carbonyl atom Oω forms a hydrogen bond with the backbone atom N of Val 54 in α GST and Leu 50 in π GST. Furthermore, in comparison to the interaction at the γ-glutamate of glutathione, the amino acid residues that interact with the terminal carboxylate atoms of glutathione glycine are dissimilar in their composition and orientation in the 4 GSTs. The Gin residue interacts with these carboxylate atoms in α GST and π GST in addition to the Arg, Lys, and Trp residues.

The orientation and alignment of glutathione thus enables the close association of the ring OH atom of residue Tyr 6 with the Sy atom of glutathione in all 4 GSTs. A notable conformational difference of this residue between Sj GST and μ GST after the superposition of the binding site is the angle of ring planes. Relative to Sj GST, this angle is −34° (+ is counterclockwise, looking away from the backbone atoms); the angle is −15° for α GST and −10° for π GST. It has been suggested that in μ GST,
the side chain of Thr 13 that sits on the Tyr 6 ring enhances the hydrogen bonding effect and the pKₐ of the glutathione cysteine (Liu et al., 1993). Valine is the corresponding residue in α GST, methionine in γ GST, and cysteine in π GST.

Site-directed mutagenesis studies have revealed the importance of specific residues in the glutathione binding site. Manoharan and coworkers (1992) showed that the substitutions of binding site residues Arg 13, Gln 62, and Asp 96 in γ GST resulted in 20–50-fold decreases of both the catalysis and glutathione binding in comparison to the wild type. However, the substitution of Tyr 7 to phenylalanine still resulted in 27% of the wild-type capacity to bind glutathione, but the enzymatic catalysis was reduced to less than 1%. The substantial reduction in the catalytic activity due to the substitution of tyrosines was also demonstrated in α GST (Liu et al., 1992). Furthermore, there are differing views on how the thiol anion of glutathione is created to promote high nucleophilic reactivity, i.e., either the thiol of glutathione is deprotonated (Liu et al., 1992) or tyrosine OH group is deprotonated (Karshikoff et al., 1993). Moreover, the importance of binding of carboxylate anions of glutathione was also shown by the mutation substitution at Trp 38 in π GST, which decreased the glutathione binding activity (Nishihira et al. 1993). Nishihira and coworkers (1993) also showed that the hydrophobic ligand binding that may occur away from the glutathione binding site caused the quenching of Trp 38 fluorescence, which suggested a conformational change of the enzyme.

Although Sj GST is from an invertebrate species and μ GST from a mammal, the 2 proteins show remarkable similarities in the amino acid sequence and structure. The shared characteristics in the 3-dimensional structures of Sj GST and μ GST extend to the similarity in biochemical properties. Walker and coworkers (1993) compared the characteristics of Sj GST (26 kDa) and Schistosoma mansoni (Sm) GST (28 kDa); whereas the latter showed catalytic properties similar to the mammalian μ class GSTs, the former gave hybridized enzymatic properties of μ and α GSTs.

In the GST binding site, glutathione binds to the domain 1 side, and there is more space on the domain 2 side for additional binding by an electrophilic substrate. This region may be called hydrophobic (H-) subsite (Reinemier et al., 1991; Garcia-Sáez et al., 1994) because the residues in this region are generally hydrophobic, but notable differences are evident. For example, in μ GST the residues Val 9, Leu 12, Ile 111, Tyr 115, Phe 208, and Ser 209 were shown to be important for the binding of 1-(S-glutathionyl)-2,4,6-trinitrocyclohexadienyl (GTD) (Ji et al., 1993) and the diastereomers of 9-(S-glutathionyl)-10-hydroxy-9,10-dihydrophenanthrene (Ji et al., 1994). The corresponding residues in Sj GST are Ile 9, Leu 12, Ser 106, Tyr 110, Gln 203, and Gly 204 (Kinemage 1). Here the key difference is the change of Phe 208 in μ GST to Gin 203 in Sj GST, for the latter is observed to interact with Arg 102. The side chain of Arg 102 point toward the binding site and glutathione. The side chain of corresponding residue Arg 107 in α GST points away from the binding site and interacts with Asp 161 that is further away. The combination of Arg 102 and Gin 203 in Sj GST could make the binding of such a molecule as GTD even more specific in Sj GST.

In the α GST structure, the ring portion of the substrate S-benzyl-glutathione is surrounded by hydrophobic residues (Val 11, Leu 106, Leu 107, Ala 215, Phe 219, and Phe 221). The binding site of α GST is less open than that of the other GSTs due to its C-terminal α-helix. In π GST, Phe 8, Pro 9, Val 10, Ala 102, and Tyr 106 may be the possible residues for xenobiotic binding interaction. The recent structure of mouse α GST (García-Sáez et al., 1994) complexed with S-nitrobenzyl-glutathione showed the specificity of these and other residues. Furthermore, Tyr 115 in μ GST is shown to be an important residue for catalytic activity (Johnson et al., 1993). The corresponding residue in Sj GST is Tyr 110, Tyr 105 in π GST, and Val 110 in α GST. The influence of this residue on catalysis in GSTs other than μ GST has yet to be determined.

Fusion peptide and antibody reactivity to Sj GST

The fusion peptide of gp41 forms a series of tight turns (Fig. 8; Kinemage 1). Muster and coworkers (1993) showed that this peptide fused to Sj GST reacted with the Mab 2F5, which strongly suggests a recognizable conformation in solution. Compared to a free peptide, the fusion peptide may be stabilized by favorable interactions with the surface of the GST protein. Computer modeling predicts that the sequence Glu-Leu-Asp-Lys-Trp-Ala is a part of an α-helix (Gallaher et al., 1989). Because only this short sequence was utilized in the fusion with Sj GST, the predicted structure may not be expected in the present crystal lattice. Moreover, a recent structure of gp120 peptide in an antibody-antigen complex also adopts a conformation of tight turns (Ghiara et al., 1994). Once determined, the larger context of

Table 2. Superposition of domain 2 after the alignment of domain 1 for the 4 GST structures*: RMS deviation, rotation angle, and magnitude of translation

<table>
<thead>
<tr>
<th>Fixed GST</th>
<th>Rotating GST</th>
<th>μ</th>
<th>α</th>
<th>π</th>
</tr>
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<tbody>
<tr>
<td>Sj</td>
<td>1.66 Å, 5.5°, 1.10 Å</td>
<td>1.43 Å, 6.8°, 3.45 Å</td>
<td>1.42 Å, 5.1°, 1.27 Å</td>
<td></td>
</tr>
<tr>
<td>μ</td>
<td>1.91 Å, 7.2°, 4.52 Å</td>
<td>1.25 Å, 4.9°, 1.32 Å</td>
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<td>α</td>
<td>1.64 Å, 8.5°, 3.98 Å</td>
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</table>

*With the amino acid sequences aligned, the Ca atoms of domain 1 (residues 2–6, 14–22, 27–32, 56–59, 62–65, and 67–77 of Sj GST) were all superimposed between different pair combinations of GSTs.
Fig. 10. Glutathione substrate (in thick lines) in the GST binding site. A: Glutathione in Sj GST. B: Glutathione in μ GST. C: S-benzyl-glutathione in α GST. D: Glutathione sulfonate in π GST.
The crystals have Laue symmetry of 4/mmm and belong to the space group P4212 with cell dimensions a = b = 94.7 Å, and c = 58.1 Å. There is 1 GST monomer per asymmetric unit. X-ray diffraction data were collected with Rigaku RAXIS-II Imaging Plate Diffractometer mounted on Rigaku RU200 X-ray Generator. The crystal to detector distance was set at 133.4 mm and generator settings were 40 kV and 70 mA. The data set contained 8,485 unique reflections with I > 1σ(I) to 2.50 Å (86.9% complete), and 10,057 reflections with I > 3σ(I) to 2.25 Å (75.9% complete). Reflections in the 2.50–2.25 Å range were recorded at the corners of the imaging plates and consequently only 45.1% complete. The Rmerge value was 11.7% for all observations with I > 1σ(I).

Molecular replacement

The structure of Sj GST was determined by molecular replacement (Rossmann, 1990) using the unmodified coordinates of the subunit A of the previously determined μ GST (Ji et al., 1992) as the starting model. The program package MERLOT (Fitzgerald, 1988) was used in these calculations. The rotation function gave 2 different solutions at different resolution limits. Both of these 2 orientations were used to calculate translation functions, but only one gave consistent peaks at various Harker sections. At the same time, the space group ambiguity was resolved. This solution was also tested by program RTMAP (written by J.X. Ho, unpubl.) that maps crystallographic R factors and correlation coefficients over all possible translation vectors for a given range of orientation angles. The translation vector that corresponded to the best R factor (47.9%) and highest correlation coefficient (30.2%) was consistent with the results from Patterson translation function for 2,248 reflections with Fc > 3σ(Fc) in the resolution range of 20.0–4.0 Å. This model was then used in a rigid body refinement after which the R factor was 30.4% and correlation coefficient 48.6% for the same group of data. The crystal packing of the molecules produces a perfect dimer of GST, i.e., 2 monomers relate to each other by a crystallographic 2-fold rotation symmetry.

Structure refinement

After the rotation and translation search, the secondary structure elements of the GST model were adjusted in the rigid body refinement performed with XPLOR program (Brünger, 1991). Then initial side-chain changes were made after alignments of the amino acid sequences of Sj GST and μ GST (Fig. 3). Direct changes were made for side chains similar in size and polarity such as Ile and Leu, and Met and Lys; other residues with pairings that did not share such characteristics were converted into alamines to produce the initial Sj GST model. Next, a simulated annealing calculation was performed with XPLOR using reflections in the 8.0–3.0-Å resolution range. Omit maps were calculated for appropriate replacements of side chains and adjustments of their geometries. Molecular graphics programs STARS (Lim, 1994) and O (Jones et al., 1991) were used to make these changes. Later, the resolution range was altered to include the 6.2-3.Å data. As the refinement progressed, water oxygen atoms were assigned for electron density peaks greater than 3σ on the Fo −Fc contour maps. The present Sj GST structure should be defined as one determined to 2.5 Å resolution; although the diffraction data extended beyond 2.5 Å resolution,
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less than 50% of the data were present in the higher resolution range. The final refinement of the Sj GST structure was performed with the restrained least-squares refinement program GPRLSA (Hendrickson & Konnert, 1980; Furey et al., 1982; Hendrickson, 1985). To determine the structure of fused gp41 peptide, the main structure of Sj GST was held fixed and the C-terminal residues were allowed to move in the simulated annealing calculations.

Determination of the main Sj GST structure

The Sj GST and μ GST proteins share 42% amino acid pair identities and many other homologous pairs (Fig. 3). One notable difference observed from the amino acid alignment is that Sj GST does not have a long loop that is present in μ GST (residues 33–42) in domain 1. Initial rigid body refinements performed without this loop resulted in the translation of as much as 1 Å for some of the α-helices, indicating that the tertiary structures of the 2 domains of Sj GST were different from those of μ GST. Furthermore, the 78–84 loop connecting the α-helices 3 and 4 was modified a few times before it was stabilized and showed a different spatial placement with respect to the 2 neighboring α-helices than the corresponding 83–89 loop in μ GST. Subtle structural differences were also noted, as when residue 154 was not changed from isoleucine of μ GST into alanine by oversight during the first several rounds of refinement. Omit map for this residue position did not fit well for isoleucine.

During the initial stages of refinement, glutathione was not included in the model. However, Fo - Fc electron density map clearly indicated its presence in the binding site (Fig. 5A). Glutathione was added, along with additional water molecules where appropriate. The presence of oxidized glutathione (2 glutathiones joined by a disulfide bond) was ruled out. As higher resolution data were included in the refinement, i.e., reflections in the 6.0–2.3-Å range, additional water molecules were gradually located.

Difficulty in the Sj GST structure refinement was encountered with the positioning of the C-terminal residues. The sequence alignment indicated that the 3 glycine residues 210–212 may be paired with the last 3 residues of μ GST (215–217), but this left the additional 5 residues of Sj GST (His 213, Asp 214, Pro 215, Pro 216, and Lys 217) and another 15 residues of the fusion peptide without a starting model. This question was resolved by the refinement performed (using reflections in the 8.0–3.0-Å range) without the residues beyond the α-helix 8. The C-terminus of Sj GST was gradually rebuilt and showed the same loop folding pattern as that for the μ GST. The presence of electron densities for 3 glycines (210–212) supported for this folding pathway. The Fo - Fc density in Sj GST could be traced to residue Pro 215 without addition of water molecules. A similar difficulty was also encountered for the structure of μ GST (Ji et al., 1992), for which the interpretation of the last 17 residues was resolved when 150 water molecules were added.

Determination of the fusion peptide structure

After the Sj GST structure was determined to residue Pro 215, the refinement was extended to higher resolution reflections in the 6.0–2.3-Å range. With the addition of approximately 50 water oxygen atoms, the location of the fusion peptide structure became more evident. Its electron density was located in the pocket formed by the α-helices 6, 7, and 8, and near Pro 215, but it was not connected to Pro 215. Examination of the crystal symmetry indicated 2 possibilities: (1) the fusion peptide extends from Pro 215 directly to this density so that it lies on the surface of the main Sj GST structure, or (2) it extends to the density in the neighboring region so that the last several residues of the fusion peptide would interact with symmetry-related GSTs other than the original Sj GST in the main cell (Fig. 7). The second possibility was a more appropriate choice, because when an alanine model was placed in the fusion peptide region, the extension of electron density was evident from the neighboring region to Pro 215. In contrast, the direct connection of the electron density on the surface of Sj GST to Pro 215 could not be made even after many rounds of model building and refinement. Following this determination, residues were added at the C-terminus to Arg 223. At this point in the sequence, the fusion peptide was now interacting with the symmetry-related GSTs. The Fo - Fc map indicated that the last 9 or 10 residues of the fusion peptide formed a loop structure. The residue Gly 224 initiates a turn, and at Met 226 another sharp turn is made. Finally, there is another turn at Asp 229 and Lys 230 (Fig. 8).

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