Rationalization of allosteric pathway in *Thermus* sp. GH5 methylglyoxal synthase

Shekufeh Zareian1, Khosro Khajeh1,*, Mohammad Pazhang2 & Bijan Ranjbar3

1Department of Biochemistry, Faculty of Biological Sciences, Tarbiat Modares University, Tehran, 2Department of Cellular and Molecular Biology, Faculty of Science, Azarbaijan University of Tarbiat Moallem, Tabriz, 3Department of Biophysics, Faculty of Biological Sciences, Tarbiat Modares University, Tehran, Iran

A sequence of 10 amino acids at the C-terminus region of methylglyoxal synthase from *Escherichia coli* (MGS) provides an arginine, which plays a crucial role in forming a salt bridge with a proximal aspartate residue in the neighboring subunit, consequently transferring the allosteric signal between subunits. In order to verify the role of arginine, the gene encoding MGS from a thermophile species, *Thermus* sp. GH5 (TMGS) lacking this arginine was cloned with an additional 30 bp sequence at the 3’-end and then expressed in form of a fusion TMGS with a 10 residual segment at the C-terminus (TMGS*). The resulting recombinant enzyme showed a significant increase in cooperativity towards phosphate, reflected by a change in the Hill coefficient (nH) from 1.5 to 1.99. Experiments including site directed mutagenesis for Asp-10 in TMGS and TMGS*, two dimensional structural survey, fluorescence and irreversible thermostability were carried out to confirm this pathway.

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INTRODUCTION

Allosteric is the regulation of the protein function and structure and/or flexibility induced by the binding of a ligand or another protein (effector) to a site which is generally far from the active site of the protein. Therefore, allostericity is the coupling of conformational changes between two separated sites. By this description, allosteric is an intrinsic property of all proteins. Effectors that elevate the protein function including ligand affinity or catalytic rate are named allosteric activators, while those that lower the function are known as allosteric inhibitors. Previously, allosteric and cooperativity were limited to multi-meric proteins but today it has also been recognized as a property of monomeric proteins (1).

Methylglyoxal synthase (MGS) (EC 4.2.3.3) catalyzes an elimination reaction that converts dihydroxyacetone phosphate (DHAP) to phosphate and pyruvaldehyde, which subsequently tautomerizes to form methylglyoxal (MG) (2, 3), in the first step of the methylglyoxal bypass of the glycolysis pathway (4-6). In the absence or presence of phosphate, MGS kinetic data are hyperbolic or sigmoidal depending on the substrate concentration, respectively (7, 8). Therefore, phosphate acts as an allosteric inhibitor (conformational) and not a product inhibitor (7). The MGS enzyme has been studied in many different organisms (9-13). However, *Escherichia coli* (E. coli) MGS is the most extensively studied enzyme (6, 8, 14-16).

Given the importance of this enzyme for scientific applications, it has been well explored; however, biological function of this substance in the metabolic pathway is still not very well known. MG is tightly bound to glycolysis from an evolutionary perspective. Although it is not in the main stream of glycolytic pathway but a role can be assigned to its production in phosphate deficiency conditions, as phosphate is one product of the related reaction (4, 17). It has been suggested that this enzyme in *E. coli* and other microorganisms may facilitate the transition between starvation and abundance (18). MG is converted either to D-lactate via the glyoxalase system or to 1,2-propandiol (a commercial commodity with industrial significance) by glycerol dehydrogenase and aldehyde reductase (4, 19, 20).

Previous studies have revealed different pathways for transition of allosteric signals from ligand to substrate binding and active sites. According to the crystallographic structure of *E. coli* MGS and mutagenic studies, two mechanisms have been proposed for transition of allosteric signals between subunits. According to the first mechanism, formation of a salt bridge between Asp-20 and Arg-150 in the presence of phosphate passes information between the six adjacent subunits. In the second putative mechanism, Pro-92, Arg-107, and Val-111 are passes information between the six adjacent subunits. In the absence or presence of phosphate, MGS kinetic data are hyperbolic or sigmoidal depending on the substrate concentration, respectively (7, 8). Therefore, phosphate acts as an allosteric inhibitor (conformational) and not a product inhibitor (7). The MGS enzyme has been studied in many different organisms (9-13). However, *Escherichia coli* (E. coli) MGS is the most extensively studied enzyme (6, 8, 14-16).

Here, we have studied MGS from a thermophile species, *Thermus* sp. GH5 (TMGS) and its allosteric regulation has been compared with *E. coli* MGS (EMGS). In the previous report (21), we analyzed the oligomerization of TMGS by size
exclusion chromatography, which showed the active fraction was related to the hexameric state (21). On the other hand, the crystallographic state of this enzyme shows the hexameric form (PDBID: 2X8W). In addition, TMGS has Hill coefficient (nH) of 1.5 in the presence of phosphate (21). Since TMGS has a lower nH compared to EMGS (7, 8), we focused on their structures at the C-terminus and assumed that the absence of the essential arginine at this terminal and consequently lack of salt bridge can be the main reason for the lower response to the transition of allosteric effects. Therefore, a segment contains 10 residues was added to TMGS C-terminus and its impact on salt bridge formation and nH was investigated. In order to confirm the role of the arginine in salt bridge formation, two additional variants, with alteration at an aspartate residue were designed. Structural features of these protein variants were also studied using circular dichroism (CD) and fluorescence spectroscopy.

RESULTS

Addition of C-terminal tail to TMGS elevated the Hill coefficient

Multiple amino acid sequence alignment of TMGS (132 residues) and E. coli MGS, a well-studied MGS with 152 residues, revealed that these two enzymes can show significant structural differences due to the loss of 20 residues in TMGS, especially the 10 amino acids that are essential for the configuration of the C-terminus α-helix in E. coli MGS (Supplementary data, Fig. S1). Saadat and Harrison have previously introduced two allosteric signal transition pathways in the E. coli enzyme, one of which happens through introduction of a salt bridge between Asp-20 and Arg-150 (16). This pathway is missing in the TMGS variant due to the loss of 10 amino acids as well as the lack of the necessary arginine at the C-terminus end. On the other hand, the comparison between the nH of TMGS (1.5) (21) and EMGS (3.4) shows a clear difference in the ability of these two enzymes in transition of the allosteric signals. However, many reasons can cause such difference, such as complete or partial cooperativity that is propounded in allosteric enzymes. It is assumed that the lack of a salt bridge in TMGS could be one of the reasons for lower allosteric response. Therefore, the question is whether the addition of an extra peptide to the C-terminus can raise the sensitivity of the enzyme to phosphate by engaging the proximal aspartate in the salt bridge.

Therefore, by designing a 60-nucleotide reverse primer, a ten amino acid sequence (YQRYLADRLK), was appended to the C-terminus of TMGS, and the addition was confirmed by gene sequencing. The new enzyme with 142 amino acid residues was called TMGS+. Moreover, the effect of this segment gene sequencing. The new enzyme with 142 amino acid residues was called TMGS+. Furthermore, SDS-PAGE analysis revealed the distinction between the mass of TMGS (14.3 kDa) and TMGS+ (15.6 kDa) as predicted (Supplementary data, Fig. S2).

Kinetic parameters of TMGS+ and TMGS were calculated from the direct assay of MGS, described in the methods section, using DHAP as a substrate and inorganic phosphate as an allosteric inhibitor. nH was raised from 1.5 in TMGS to 1.99 in TMGS+ which revealed the effect of arginine at position 140. Modifications were designed in both TMGS+ and TMGS to assess the salt bridge arrangement between Arg-140 and the proximal aspartate at position 10 in TMGS (equal to Asp-20 in EMGS).

Mutation of Asp-10 to Asn confirmed the salt bridge arrangement in TMGS+

TMGSD10N and TMGS+D10N mutants were constructed, using the quick change method. Cloning, expression and purification of enzymes were done as described above. The kinetic parameters and Hill coefficients were also calculated. Comparison of these parameters in four variants could reveal the effect of the embedded Arg-140 and subsequent formation of a salt bridge between this arginine and Asp-10.

In spite of an increase in TMGS+ nH from 1.5 to 1.99, this coefficient did not raise in TMGS+D10N despite the presence of a C-terminal Arg-140. Furthermore, the TMGSD10N variant did not show any sensitivity to this mutation. Km, kcat and catalytic efficiency of the mutants and their nH, in 0 and 1.5 mM phosphate are presented in Table 1. In addition, the Michaelis-Menten curve and Hill plot of four variants are depicted in Fig. 1. The formation of the helical structure as a result of tail attachment to TMGS was investigated by secondary structure analysis using circular dichroism spectra measurements.

Circular dichroism spectra of TMGS and TMGS+

The Segment added to the TMGS C-terminus was assumed to form a helical structure like its equivalent part in EMGS. Structural studies were performed to confirm the presence of an α-helical structure as an efficient spatial structure for arrangement of a salt bridge. As seen in Fig. 2 helical structure of the TMGS+ was increased compared to TMGS. Further, the Enzymes compactness and thermal stability were measured by intrinsic fluorescence and irreversible thermonactivation assay.

Table 1. Kinetic parameters of TMGS, TMGS+ and two mutated enzymes

|          | Km (mM) | kcat (s⁻¹) | kcat/Km | nH⁺   |
|----------|---------|------------|---------|-------|
| Thermus sp. GHS |         |            |         |       |
| MGS (TMGS) | 0.50 ± 0.02 | 326.4 ± 2 | 652.9   | 1.50  |
| TMGS+      | 0.21 ± 0.01 | 102.9 ± 1  | 490.0   | 1.99  |
| TMGS+D10N  | 0.75 ± 0.03 | 0.41 ± 0.02 | 5.44    | 1.40  |
| TMGS*D10N  | 0.25 ± 0.01 | 0.09 ± 0.004 | 0.374  | 1.44  |

*Hill coefficient is for 1.5 mM concentration of phosphate.
Fig. 1. Michaelis-Menten curve and Hill plot of four variants in different concentrations of DHAP as substrate. (A) 0 mM phosphate, (B) 1.5 mM phosphate (C) Hill plot of four variants. Hill plot was drawn according to the equation mentioned in the Materials and Methods section, in the presence of 1.5 mM of phosphate. (♦) TMGS, (▲) TMGS D10N, (■) TMGS+, (●) TMGS+D10N.

Fig. 2. Far-UV CD spectra of TMGS (---) and TMGS+ (--). Diagram reveals a more helical structure for TMGS+.

Intrinsic fluorescence
It is assumed that TMGS+ has a more compact structure than TMGS due to the formation of a salt bridge in its structure. The fluorescence emission of a protein is the result of its intrinsic fluorophores such as tryptophan and tyrosine residues. The emission spectrum of these aromatic residues is highly sensitive to their surrounding environment. Structural changes in protein may expose the internal tryptophan residues to aqueous environment or bury the accessible residues in the core of the protein. Alterations in the polarity of tryptophan microenvironment will change the emission spectrum, therefore, it is possible to indirectly follow the conformational changes of the enzyme through its intrinsic fluorescence emission. Upon excitation at 280 and 293 nm, TMGS and TMGS+ showed maximum emission intensity at 340 nm but TMGS+ demonstrated a much higher intensity. Since more structural compactness leads to a higher intensity we assume that a salt bridge might have formed between two adjacent subunits (Asp-10 and Arg-140) that may bury some of the fluorophores in a more hydrophobic microenvironment (Fig. 3).

Irreversible thermoinactivation study of TMGS and TMGS+ to compare structural stability
Thermal inactivation of TMGS+ and TMGS was performed in support of the fluorescence spectroscopy data. The results show that TMGS+ with a half life of 10 min was more stable than TMGS with a half life of 3 min, at 80°C, which suggests a more compact and stable structure in TMGS+. The results are depicted in Fig. S3.

DISCUSSION
The aim of this study was to explore a phenomenon observed by our group (21), which revealed that methylglyoxal synthase from Thermus sp. GHS (TMGS) has a lower nH (1.3) com-
pared to MGS from *Escherichia coli* (EMGS) (3.4). The experiments aimed to identify the reason for lower sensitivity to phosphate concentration and consequently weaker transmission of allosteric signals between neighboring subunits in TMGS. In a study by Saadat and Harrison, two pathways of transmitting the allosteric effect have been proposed for the *E. coli* MGS. In the first possible mechanism, formation of a salt bridge between Asp-20 and Arg-150 was proposed in presence of phosphate that passes on the information between the six adjacent subunits, while in the second proposed mechanism, Pro-92, Arg-107, and Val-111 are employed in this convey. The three amino acids involved in the second pathway are highly conserved among different strains; however, the first pathway has been lost in TMGS due to the absence of the essential arginine which participates in salt bridge formation between subunits. Therefore, to investigate the effect of salt bridge formation on cooperativity we decided to add a segment (10 residues) containing the mentioned arginine to the C-terminus end. An increase in $nH$ of TMGS has not increased as much as $nH$ of EMGS (3.4). To explain high cooperativity of EMGS further investigations are required.

Overall, in this work, importance of salt bridge formation between Arg-140 and Asp-10 in the creation of cooperative TMGS has been investigated which shows that this pathway could increase $nH$ of TMGS from 1.5 to 1.99 but this value is lower than $nH$ of EMGS (3.4). The CD spectra demonstrated a more helical structure like its equivalent part in EMGS was tested by CD analysis. The CD spectra demonstrated a more helical structure in TMGS$^+$ in comparison with TMGS. However, this data is not an evidence of structural compactness due to formation of salt bridge between Arg-140 and Asp-10 in TMGS$^+$. Fluorescence spectroscopy data demonstrated a more compact structure for TMGS$^+$ compared to TMGS, which could be explained by a salt bridge formation between adjacent subunits. Addition of two tyrosines to TMGS$^+$ tail could be the main cause of higher fluorescence intensity at 340 nm when excited at 280 nm; therefore, the samples were also excited at 293 nm, which specially reveals the tryptophan emission in both enzymes. On the other hand, TMGS$^+$ shows a higher thermal stability at 80°C.

Finally, these experiments may untie a problem in the big hasp of protein science but still the main question remains; why the mesophilic MGS (EMGS) and thermophilic MGS (TMGS) show different responses to the phosphate concentrations and consequently produce different amounts of methylglyoxal. This ability may have been acquired (through evolution) by *E. coli* in the periods of time.

**MATERIALS AND METHODS**

**Chemicals**

T4-DNA ligase and restriction enzymes were purchased from Fermentas (Vilnius, Lithuania). Oligonucleotides were synthesized by MWG Company (Germany). Tryptone and yeast extract were from Liofilchem (Roseto degli Abruzzi, Italy). Dihydroxyacetone phosphate was purchased from Sigma-Aldrich (USA). 2,4-Dinitrophenylhydrazine and other chemicals were obtained from Merck (Darmstadt, Germany).

**Construction of TMGS$^+$ plasmid**

The TMGS$^+$ gene, (TMGS with additional 30 bp at 3′-end) was constructed using pET-21a plasmid containing the TMGS gene as a template (21). 0.08 μM forward (5′-GGAATTCCATATGCGAGCGCTCCGGCGGATTG-3′) and reverse (5′-GCGAATACCGGTTATATGCGCCTGCATTG-3′) primers were used to amplify TMGS$^+$, in the presence of 1.25 unit taq-polymerase (with an...
annealing temperature of 72°C using polymerase chain reaction thermocycler. The resulting fragment (429 bp) digested with *hindIII* and *NdeI* (the underlined bases) and ligated into the similarly digested pET-21a(+) using T4-DNA ligase. The accuracy of final product was confirmed by DNA sequencing.

**Mutagenesis**

Site directed mutagenesis was carried out, as described by Fisher and colleagues, using Quick-Change method and chemically synthesized oligonucleotide primers (Bioneer, South Korea) (22). Plasmid pET-21a (+) containing the TMGS gene was used as the template. PCR reaction was carried out in the 50 μl volume containing DNA template (10 ng), 10 × PCR buffer, 0.2 mM of each dNTP, 0.8 μM of each primer and PWO polymerase (1.25 units). The mixture was heated at 95°C for five minutes and then subjected to 22 cycles of thermal cycling at 94°C for 1 min, 55°C for 1 min and 68°C for 13 min. PCR product was incubated with DpnI at 37°C for 12 h and then transformed to *E. coli* XL1-blue (23). Five clones were randomly selected for sequencing and confirmation of the mutations.

In order to construct TMGS*D10N* gene, TMGS*D10N* gene was used as template and the other steps were carried out as described previously.

**Expression and purification of the enzyme**

Constructed expression plasmids were transformed into the *E. coli* strain BL21 (DE3), to produce the 15.6 kDa methylglyoxal synthase and its variants. The bacteria containing recombinant plasmids were grown in Luria-Bertani (LB) medium supplemented with ampicillin (100 μg/ml) at 37°C, 220 rpm. IPTG (1 mM) was added to the culture medium once the culture reached an optical density of 0.5 to 0.7 at 600 nm. After 19 h, cells were harvested by centrifugation at 5,000 rpm for 20 min and resuspended in lysis buffer containing 2 mM imidazole, 1 M NaCl and 50 mM Tris (pH 7.0). The suspension was subjected to sonication and total lysate was centrifuged for 20 min at 12,000 rpm at 4°C. The supernatant underwent heat-shock at 70°C for 15 min, and the precipitated proteins were removed by centrifugation at 14,000 rpm for 20 min at 4°C (freezing followed by heating the supernatant improved the purification efficiency). The resulted supernatant was centrifuged at 50,000 rpm for 20 min at 4°C (freezing followed by heating the supernatant improved the purification efficiency). The supernatant were incubated at 60°C for 5 min. Then the methyglyoxal concentration (7, 14, 26).

**Enzyme kinetics**

Kinetic parameters were determined as explained above. Mixtures containing different concentrations of DHAP with no enzyme were used as controls and each data point (initial velocity) was determined in triplicate. Steady-state kinetic parameters in the presence and absence of phosphate were fitted to Michaelis-Menten equation and were numerically analyzed by Lineweaver-Burk equation. nH was calculated from the following equation:

\[
\log\left(\frac{v}{V_{\text{max}} - v}\right) = nH \log[S] - \log(K')
\]

Where \(v\) and \(V_{\text{max}}\) are velocity and maximal velocity of the enzyme, respectively, and \(nH\) is the Hill coefficient. \(K'\) is related to \(K_m\) but also contains terms related to the effect of substrate occupancy at one site on the substrate affinity of the other sites. According to this equation, the value of \(nH\) can be calculated by plotting \(\log[v/(V_{\text{max}} - v)]\) against \(\log[S]\).

**Fluorescence measurements**

Tryptophan fluorescence of TMGS and TMGS*D was measured using a Perkin Elmer luminescence spectrometer LS 55. Samples were excited at 295 nm and the emission was recorded between 300 to 400 nm. All experiments were carried out at room temperature and protein concentrations were 20 μM in 20 mM Tris buffer (pH 8.0).

**Irreversible thermoinactivation**

Thermal inactivation of TMGS and TMGS*D was investigated in 50 mM imidazole buffer (pH 6.0) at 80°C. Periodically, Samples were removed, cooled on ice and then residual activity was determined as described above. The untreated sample was used as a control (100% activity).

**Circular Dichroism measurements**

Far-UV spectra (190-260 nm) were recorded on a Jasco spectropolarimeter J-715 (Tokyo, Japan) using 1 mm path length...
quartz cell at the protein concentration of 0.2 mg/ml in 20 mM tris buffer (pH 8.0). Results are presented as molar ellipticity \( [\theta] \) (deg cm\(^2\) dmol\(^{-1}\)) based on a mean amino acid residue weight of 110 for TMGS and TMGS\(^{-}\). The molar ellipticity \( [\theta] \) was calculated from the formula \( [\theta] = \theta \times 100 \text{ MWR} / l \), where \( c \) is the protein concentration in mg/ml, \( l \) the light path length in centimeters, and \( \theta \) is the measured ellipticity in degrees wavelength \( \lambda \). The secondary structure parameters were calculated using J700 CD-JASCO software.

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