Thermostability of α-Galactosidase AgaB1 in *Thermus thermophilus*

Olafur Fridjonsson,* Hildegard Watzlawick, and Ralf Mattes

Institut für Industrielle Genetik, Universität Stuttgart, 70569 Stuttgart, Germany

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The evolutionary potential of a thermostable α-galactosidase, with regard to improved catalytic activity at high temperatures, was investigated by employing an in vivo selection system based on thermophilic bacteria. For this purpose, hybrid α-galactosidase genes of *agaA* and *agaB* from *Bacillus stearothermophilus* KVE39, designated *agaA1* and *agaB1*, were cloned into an autonomously replicating *Thermus* vector and introduced into *Thermus thermophilus* OF1053GD (*ΔagaT*) by transformation. This selection strain is unable to metabolize melibiose (α-galactoside) without recombinant α-galactosidases, because the native α-galactosidase gene, *agaT*, has been deleted. Growth conditions were established under which the strain was able to utilize melibiose as a single carbohydrate source when harboring a plasmid-encoded *agaA1* gene but unable when harboring a plasmid-encoded *agaB1* gene. With incubation of the *agaB1* plasmid-harboring strain under selective pressure at a restrictive temperature (67°C) in a minimal melibiose medium, spontaneous mutants as well as *N*-methyl-*N*-nitro-*N*-nitrosoguanidine-induced mutants able to grow on the selective medium were isolated. The mutant α-galactosidase genes were amplified by PCR, cloned in *Escherichia coli*, and sequenced. A single-base substitution that replaces glutamic acid residue 355 with glycine or valine was found in the mutant *agaB1* genes. The mutant enzymes displayed the optimum hydrolyzing activity at higher temperatures together with improved catalytic capacity compared to the wild-type enzyme and furthermore showed an enhanced thermal stability.

To our knowledge, this is the first report of an in vivo evolution of glycoside-hydrolyzing enzyme and selection within a thermophilic host cell.

Stability of enzymes and activity at high temperatures are important and desirable properties for various biotechnological processes. Hence, improving stability and activity of enzymes has been the major aim of many applied studies. Thereby, randomization methods for in vitro protein evolution, such as random mutagenesis for obtaining point mutations, DNA shuffling, and elongation mutagenesis, have been successfully applied (12, 18, 23, 24). Another approach for enhancing enzyme properties is to use in vivo evolution systems. Thermostable enzyme variants may be established by cloning mesophilic genes into a thermophilic organism and applying a growth selection where elevated temperatures function as a selective pressure. Such a thermostabilization was applied, e.g., by the stabilization of kanamycin nucleotidyltransferase in *Bacillus stearothermophilus* (17, 22) and by the stabilization of a 3-isopropylmalate dehydrogenase (of the leucine biosynthetic pathway) in *Thermus thermophilus* (29). Thermostabilization can also be supported by in vitro mutagenesis followed by a growth selection in a thermostable as described by Kotsuka et al. (13) for the further stabilization of 3-isopropylmalate dehydrogenase in *T. thermophilus*.

Thermostabilization of proteins may also involve an improvement of properties such as substrate affinities of enzymes at high temperatures. We have studied thermostable α-galactosidases from various mesophilic and thermophilic bacteria, including the isoenzymes *AgaA* and *AgaB* from *Bacillus stearothermophilus* KVE39 (7, 9, 11, 32). Although the amino acid sequences of *AgaA* and *AgaB* are highly similar (97% identity), the enzymes exhibit different properties (11). *AgaA* displays a high affinity for oligosaccharides such as melibiose and raffinose and a temperature optimum of activity at 65 to 67°C under given assay conditions. On the other hand, *AgaB* displays a low affinity for melibiose and raffinose and a temperature optimum of activity at 45 to 50°C. Yet *AgaB* shows a marginal stability at 60 to 65°C (11, 32). In a previous work on *AgaA* and *AgaB*, it was demonstrated that a central BsmI-HindIII fragment in their encoding genes, designated *agaA* and *agaB*, respectively, could account for the characteristic phenotypes of the enzymes, i.e., the temperature optimum of activity and the hydrolyzing rate (11, 32; H. Watzlawick, unpublished results). The aim of this work was to establish a selection system for α-galactosidases based on a growth selection of thermophilic bacteria, thereby achieving high enzyme activity and stability at elevated temperatures. The fact that the phenotypes of *AgaA* and *AgaB* are encoded in genetically exchangeable elements allowed hereby the application of the hybrid genes *agaA1* and *agaB1* as test genes for the selection system. The hybrid genes differed only in respect to their central BsmI-HindIII fragment (Fig. 1). This simplified plasmid construction protocols; besides, mutations leading to the *AgaA* phenotype from the *AgaB* phenotype were expected to take place in this central fragment.

Expression of heterologous genes and a subsequent thermostabilization of the gene products require the inactivation of analogous genes in the selector strain. A suitable host strain for a selection of thermostable α-galactosidases active at elevated temperatures was constructed by applying integration mutagenesis in combination with phenotypic selection and designated *T. thermophilus* OF1053GD, as previously described (8). In this paper we describe the cloning of the α-galactosidase gene derived from *B. stearothermophilus* in the thermophilic bacterium *T. thermophilus* and a subsequent selection of en-

* Corresponding author. Present address: Prokaria Ltd., Gylfaflöt 5, 112 Reykjavik, Iceland. Phone: (354) 570 7914; Fax: (354) 570 7901. E-mail: olafur@prokaria.com.
zyme variants, with an increased temperature optimum of activity and enhanced stability, following adaptation (isolation of spontaneous mutants) or in vitro mutagenesis.

MATERIALS AND METHODS

Bacterial strains, growth conditions, and transformation procedure. *Thermus thermophilus* OF1053GD (8) (∆*agaT* ∆*kan* Gal*), a mutant of strain TH125 (10), was used as a host strain for the cloning and the selection procedure. The strain was grown under strong aeration in mineral medium 162 (4) with 0.25% tryptone and 0.25% yeast extract at pH 7.5 (T162). Growth was carried out at 65°C under nonselective conditions and at 60°C when cultures contained kanamycin (20 mg ml⁻¹) for plasmid selection. Growth on a single carbon source (melibiose, 0.1, 0.2, and 0.4%) was tested on agar plates containing the minimal medium 162 (4) with a slight modification as previously described (8). The method of Koyma et al. (14) was used for the transformation of *T. thermophilus* with a slight modification as previously described (7). Transformants were selected on T162 agar plates containing 20 μg of kanamycin ml⁻¹ and incubated at 60°C for 48 h. All plasmids constructed, except those which could replicate only in *T. thermophilus*, were brough into *Escherichia coli* 3B109 [pCF44 Δ(lac-pro4B) bsdR17 recA1 endA1 gyrA96 thi-1 relA1] (F' traD36 proAB lacF2 ΔM15) (31) by transformation (2). Transformants were either selected for ampicillin resistance on agar plates with 100 μg of ampicillin per ml or for kanamycin resistance on plates with 25 μg of kanamycin per ml.

DNA manipulation and general plasmid construction. Recombinant DNA techniques, i.e., plasmid preparation, subcloning, and agarose gel electrophoresis, were performed by conventional protocols (25). Structures of the plasmids were confirmed by restriction mapping. Further, the inserts of pOF287 and pOF288 were confirmed by sequencing. Sequencing reactions of double-stranded DNA were carried out according to the dyeoxy chain termination method with universal and internal primers (26). The plasmids used in this study are listed in Table 1.

Construction of *Thermus* replication plasmids containing *agaT* and *agaB* hybrid genes. Hybrid genes, based on *agaA* and *agaB*, of *B. stearothermophilus* KVE9 were cloned into a *Thermus* autonomously replicating plasmid, downstream of the E. coli tac promoter (1) and a ribosome binding site from *T. thermophilus* (8). The construction of plasmid pOF2811, carrying an AgaA type encoding gene, *agaA1*, which served as a positive control, was as follows. Plasmid pOF1253 contains *agaA1* downstream of the *Thermus* ribosome binding site. It was constructed by replacing the *kan* gene of pOF1155 (8), which contains the marker between flanking sequences of the native *o-galactosidase gene* *agaT* (7), with *agaA1*. Due to an NdeI restriction site in *agaA*, a three-frame ligation was performed with the following: the *agaA* gene following amplification (8) and *Nde*-HindIII digestion; *agaB*, following amplification (8) and HindIII-BglII digestion; and the pOF1155 following *Nde*-BamHI digestion. The resulting chimeric gene encoded an enzyme with the characteristic phenotypes of AgaA. The construction of pOF10726 is described elsewhere (8). The plasmid contains a stable plasmid mutant of pOF5714 (8) in a pUC19 derivative. A PstI fragment from pOF10726 containing repA4, the minimal replication unit (3) from the *Thermus* plasmid pTSPI (16), was cloned in pC20R (19) to produce pOF184. *agaA1* along with the *Thermus* putative Shine-Dalgarno (SD) sequence from pOF1253 was amplified by PCR with primers S1065 and S952 (8). The EcoRI restriction site in the S1065 primer was disrupted by EcoRI digestion and a Klenow modification. This was done in order to be able to use other EcoRI restriction sites in later steps. Subsequently, the gene was blunt end ligated downstream of the tac promoter in pBFacI (1) to produce pOF182. Further, *agaA1* along with the tac promoter and the putative *Thermus* SD sequence was amplified from pOF182 by PCR with primers S1318 and S952 (8) and cloned between the HindIII and BglII sites of pOF184. This was carried out in two steps. First, the 5′ region of *agaA* (a HindIII-BglII fragment) was ligated between the HindIII and BglII sites of pOF184 to produce pOF185. Then, the 5′ region of *agaA*, along with the upstream P*gaB* sequence, was ligated into the HindIII site of pOF185 to produce pOF187. The *kan* gene (21), along with the *Thermus* agaA promoter (16) from pOF1056 (8), was ligated into the Muni site of pOF187 to produce pOF287. The pIC region of pOF287 was deleted by EcoRI digestion, and the remaining plasmid was self-ligated to produce pOF2811 before transformation of *Thermus thermophilus*. The corresponding AgaB type plasmid, pOF2812, was generated in the same way except the plasmid pOF762 was the initial source of *agaB*. To construct pOF762, a BsmI-MunI fragment of pOF1253 was replaced by a corresponding agaB BsmI-MunI fragment following the amplification of *agaB* from pCG3 (11), using the primers S951 and S952 (8) and a BsmI-MunI digestion and a ligation. All plasmids involved in the construction of pOF2811 and pOF2812 are listed in Table 1. Figure 2 shows the last step in the construction of pOF2812 containing *agaB1*, which was used to transform *T. thermophilus* OF1053GD for subsequent thermoadaptation experiments.

**Thermoadaptation in aequorin culture.** N-methyl-N-nitro-N-nitrosoguanidine (MNNG) plasmid mutagenesis, and reclfoning into *E. coli*. Cells from a late-exponential-phase culture of *T. thermophilus* OF1053GD harboring pOF2812, in T162 kanamycin medium. Several concentrations of MNNG were tested (0.05 to 0.25 mg ml⁻¹). Following growth overnight at 60°C, the cells were washed in 0.9% NaCl and plated on 162 minimal melibiose agar medium. Plasmids from strains, capable of growth, were isolated by alkali lysis (25) and reintroduced into a plasmid-free strain of OF1053GD by transformation for the confirmation of the plasmid-dependent phenotype. Further, the mutant genes were amplified by PCR, with primer S462, CGG AAT TCA GAG AAT GTC, following incubation for 6 days, the cells were diluted (three times) in a fresh medium. Following incubation for 13 days, the culture was diluted and plated on a minimal melibiose agar medium.

Plasmid mutagenesis was performed by adding MNNG to an exponential culture (optical density at 600 nm, 0.4) of *T. thermophilus* OF1053GD harboring pOF2812, in T162 kanamycin medium. All plasmid constructs were transferred into *T. thermophilus* by transformation. Following incubation for 6 days, the cells were diluted (three times) in a fresh medium. Following incubation for 13 days, the culture was diluted and plated on a minimal melibiose agar medium.

**TABLE 1. Plasmids used in this study**

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Description</th>
<th>Reference</th>
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<tr>
<td>pIC20R</td>
<td>Ap', PBR322 ori lacZ</td>
<td>19</td>
</tr>
<tr>
<td>pCG1</td>
<td><em>agaA</em> in pUC12</td>
<td>11</td>
</tr>
<tr>
<td>pCG3</td>
<td><em>agaB</em> in pUC12</td>
<td>11</td>
</tr>
<tr>
<td>pOF1056</td>
<td>P&lt;sup&gt;sal&lt;/sup&gt;-<em>kan</em> from PM1Y in pJOE930</td>
<td>8</td>
</tr>
<tr>
<td>pOF5714M</td>
<td><em>agaA</em> downstream of the P&lt;sup&gt;sal&lt;/sup&gt; in <em>Thermus</em> replication plasmid</td>
<td>8</td>
</tr>
<tr>
<td>pOF1176</td>
<td><em>agaB2</em> downstream of the P&lt;sup&gt;sal&lt;/sup&gt; in <em>Thermus</em> replication plasmid</td>
<td>8</td>
</tr>
<tr>
<td>pOF10726</td>
<td>pOF5714M cloned into a pUC derivative</td>
<td>8</td>
</tr>
<tr>
<td>pOF1155</td>
<td>kan downstream of the 5′ flanking sequence of <em>agaT</em> from <em>T. thermophilus</em> in a pUC derivative</td>
<td>8</td>
</tr>
<tr>
<td>pOF1253</td>
<td><em>agaA1</em> in place of <em>kan</em> in pOF1155</td>
<td>This study</td>
</tr>
<tr>
<td>pOF762</td>
<td><em>agaB1</em> in place of <em>agaA1</em> in pOF1253</td>
<td>This study</td>
</tr>
<tr>
<td>pOF182</td>
<td><em>agaA1</em> with a <em>Thermus</em> SD sequence in <em>E. coli</em></td>
<td>This study</td>
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<tr>
<td>pOF183</td>
<td><em>agaB1</em> with a <em>Thermus</em> SD sequence in <em>E. coli</em></td>
<td>This study</td>
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<td>pOF184</td>
<td>pTSPI region of pOF10726 in pIC20R</td>
<td>This study</td>
</tr>
<tr>
<td>pOF185</td>
<td>3′ region of <em>agaA1</em> in pOF184</td>
<td>This study</td>
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<td>pOF187</td>
<td><em>agaA1</em> downstream of P&lt;sup&gt;sal&lt;/sup&gt; and a <em>Th. thermophilus</em> putative SD sequence in pOF184</td>
<td>This study</td>
</tr>
<tr>
<td>pOF188</td>
<td><em>agaB1</em> downstream of P&lt;sup&gt;sal&lt;/sup&gt; and a <em>Th. thermophilus</em> putative SD sequence in pOF184</td>
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<td>pOF287</td>
<td>P&lt;sup&gt;sal&lt;/sup&gt;-<em>kan</em> module from pOF1056 in pOF187</td>
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<td>pOF288</td>
<td>P&lt;sup&gt;sal&lt;/sup&gt;-<em>kan</em> module from pOF1056 in pOF188</td>
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<td>pOF2811</td>
<td>pIC region deleted from pOF287 (agaA)</td>
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<tr>
<td>pOF2812</td>
<td>pIC region deleted from pOF288 (agaB1)</td>
<td>This study</td>
</tr>
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Enzyme assays. Preparation of crude extracts, estimation of protein concentrations, and procedure of enzyme assays have been described previously (7). α-Galactosidase activity was determined by measuring the rate of para-nitrophenyl-α-D-galactoside (pNP-α-galactoside) hydrolysis (4 mg ml⁻¹ in 0.1 M potassium phosphate buffer [pH 6.5]) at the desired temperature. One unit of activity is defined as the amount of enzyme required to hydrolyze 1 μmol of pNP-α-galactoside min⁻¹ under a given assay condition (9). Further, the rate of τ-talassone or melibiose hydrolysis, in 0.1 M potassium buffer, pH 6.5, was determined by measuring the amount of α-galactose released using high performance liquid chromatography equipment (7). Colonies, which displayed α-galactosidase activity, were detected by histochemical staining as previously described (8).

Determination of temperature optimum and stability. Production of enzymes for characterization was achieved with E. coli crude extracts (1 mg of protein ml⁻¹) at 60°C for 20 min. The enzymes were then incubated in 0.1 M potassium phosphate buffer (pH 6.5) for 30 min at various temperatures over a range from 25 to 80°C. Following incubation the enzyme was cooled, and the remaining activity was determined at the same temperature for all enzymes (37°C).

RESULTS AND DISCUSSION

Establishing a selection system. Our intention was to subject AgaB1 to thermoadaptation for selection of enzyme variants, which supported growth of the host strain on minimal melibiose medium at 67°C. For the establishment of a selection system, we intended to use AgaA1 as a positive control. In our previous work (8), we inserted the α-galactosidase genes agaA and agaB into a Thermus autonomously replicating vector downstream of the strong slpA promoter (16). Cells carrying these plasmids displayed a high-level production of α-galactosidases, verified by sodium dodecyl sulfate gel electrophoresis and activity tests (8). Although cells harboring the agaA plasmid (pOF5714) grew at 67°C on agar plates containing minimal melibiose medium, a noncontinuous growth in minimal melibiose aqueous medium was observed (data not shown). This was interpreted as an effect of the high-level α-galactosidase gene expression. In our previous work, we observed that a Gal⁷ phenotype of T. thermophilus was essential for growth on melibiose (8). As the host strain OF1053GD was capable of growing only slowly on a minimal agar medium containing 0.3% galactose, it is possible that the galactose metabolism could not fully compensate for the large amount of galactose produced from the melibiose hydrolysis in the cell, due to the high production of AgaA. Therefore, the α-galactosidase production was confined by constructing new plasmids with α-galactosidase genes inserted downstream of a Thermus ribosome binding site and the E. coli tac promoter, which was known to show low activities in T. thermophilus (20). Concomitantly, the kanamycin selection marker (21) was inserted downstream of the slpA promoter as in the shuttle vector pMY1 (16). To facilitate the construction of the plasmid modules, hybrid genes of agaA and agaB, designated agaA1 and agaB1, respectively, were used. The enzyme activities in crude extracts of T. thermophilus OF1053GD harboring pOF2811 with agaA1 or pOF2812 with agaB1 are shown in Table 2.

For the comparison, enzyme activities in T. thermophilus OF1053GD harboring pOF5714 (agaA) and pOF1176 (agaB2) from our previous

<table>
<thead>
<tr>
<th>Temp (°C)</th>
<th>pOF2811 (AgaA1)</th>
<th>pOF2812 (AgaB1)</th>
<th>pOF5714M (AgaA)</th>
<th>pOF1176 (AgaB2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>50⁰</td>
<td>1.1</td>
<td>3.4</td>
<td>12.1</td>
<td>46.8</td>
</tr>
<tr>
<td>60⁰</td>
<td>2.2</td>
<td>2.2</td>
<td>23.4</td>
<td>30.6</td>
</tr>
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</table>

*Activity tests (in triplicates) were performed using pNP-α-galactopyranoside as substrate. The maximum variation from the mean values (shown) was less than 5%.

For more details about pOF5714M and pOF1176, see reference 7.
work (8), with the α-galactosidase genes downstream of the slpA promoter (16), are shown. Purified AgaA and AgaB2 displayed specific activities similar to those of AgaA1 and AgaB1, respectively (data not shown). As expected, a roughly 10-fold lower level of enzyme activity was measured in cells containing the α-galactosidase genes downstream of the tac promoter. Moreover, the strain OF1053GD harboring the agaA1 plasmid pOF2811 exhibited stable growth in a liquid minimal melibiose medium (0.4%). In contrast, OF1053GD without a plasmid or harboring the agaB1 gene in the plasmid pOF2812 remained in the lag phase for a period of days (Fig. 3). Also, growth at 67°C on minimal agar plates containing 0.2 and 0.1% melibiose was observed only with strain OF1053GD carrying the agaA1 gene in plasmid pOF2811 (Table 3) following incubation for 5 days. Thus, the requirements for the genetic selection were achieved.

Thermoadaptation of AgaB1, MNNG mutagenesis, and screening. T. thermophilus OF1053GD harboring pOF2812 (agaB1) was incubated at 67°C in an aqueous minimal medium containing 0.4% melibiose. The cells divided once and then remained in the lag phase. Although the cells did not grow at this temperature during the first days, no death phase was observed, according to cell titer quantification (data not shown). Slow growth was observed after 11 days of incubation. The cells were plated out and tested for growth on minimal agar medium. A thermoadapted strain harboring a plasmid, designated pOF2812 M1, was selected for further examination. Subsequently, the plasmid was isolated from its host strain and designated pOF2812 M1, was selected for further examination. Plasmid mutagenesis is commonly applied by subculturing overnight cultures into fresh minimal medium, thus confirming the plasmid-dependent phenotype. Further, α-galactosidase activity at 50 and 60°C in crude extracts of the plasmid-harboring strains was examined. All enzymes displayed higher activity at 60 than at 50°C, in contrast to the wild-type enzyme AgaB1.

Sequencing and characterization of the mutant enzymes. The sequencing of the mutant genes, designated agaM1 to agaM5 (from pOF2812 M1 to M5, respectively), revealed a single nucleotide replacement in all cases. In agaM1 to agaM4, A→1064, according to the Bsm1-HindIII fragment of the gene and corresponds to a Glu355→Gly amino acid replacement. In agaM5, the same nucleotide (A-1064) was replaced with T, which corresponds to a Glu355→Val amino acid replacement. These base substitutions are inconsistent with the generally accepted effect of MNNG mutagenesis, which most commonly leads to GC→AT transitions (27). Nevertheless, control plating indicated the mutagenic effect of MNNG. No colonies were detected during the mutagen approach, on minimal agar medium following 6 days of incubation, where cells without a mutagenic treatment had been plated as a control. On the other hand, the A1064→G transition in pOF2812 M1, which occurred spontaneously, was detected following a prolonged incubation at a restricted temperature. Although MNNG-induced A→G transitions or A→T transversions are generally infrequent, such mutations induced with alkylating agents are common in E. coli alkA mutants, which are deficient for 3-methyladenine DNA glycosylase II, involved in induced DNA repair activities. In most cases these mutations are umaC dependent, i.e., resulting from SOS mutagenic processing (6). Thus, it is possible that deficient specific DNA repair activities in T. thermophilus OF1053GD, induced by exposure of MNNG, caused the base substitutions at A/T site 1064 to occur. Also, this site may be a substitution hot spot, similar to what is known to occur in E. coli (5).

The enzymatic properties of AgaM4 (Glu→Gly) and AgaM5 (Glu→Val) were examined and compared with those of the wild-type enzyme (AgaB1). Figure 4 shows the effect of temperature on the activity and stability (panels A and B, respectively). The optimum hydrolyzing activity of AgaM4 (following 15 min of preincubation at respective temperatures) was determined to be at 65°C, about 15°C higher than for the wild-type enzyme (50°C). The optimum activity of AgaM5 was determined to be at 60°C (Fig. 4A). The temperature where 50% mids, including pOF2812 M1, with the spontaneous mutation, were all capable of growing at 67°C on a minimal melibiose medium, thus confirming the plasmid-dependent phenotype. Further, α-galactosidase activity at 50 and 60°C in crude extracts of the plasmid-harboring strains was examined. All enzymes displayed higher activity at 60 than at 50°C, in contrast to the wild-type enzyme AgaB1.

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of the activity remains following a 30-min incubation (T[1/2]) was determined to be 71°C for the mutant enzymes, at least 3°C higher than for the AgaB1 enzyme (Fig. 4B). Although this difference is not large, these results were reproducible. The slightly enhanced stability observed for the mutant enzymes can, however, hardly account for the increase in the temperature optimum for the hydrolyzing activity. Table 4 summarizes the \( K_m \) values for pNP-\( \alpha \)-galactosidase hydrolysis at 25, 50, and 60°C. The \( K_m \) of AgaB1 increases substantially along with a rise in temperature, whereas the \( K_m \) values of the mutant enzymes remain below 0.5 mM. Thus, the hydrolyzing rate of AgaB1 at high temperatures is affected by the drastic change in substrate affinity. Table 5 summarizes the kinetic parameters of AgaA, AgaB1, and the mutant enzymes for melibiose and raffinose hydrolysis. The \( K_m \) values for AgaM4 and AgaM5 are much lower than for AgaB1 and resemble the values of AgaA. Further, the mutant enzymes are superior to the progenitor enzyme (AgaB1) in regard to catalytic properties and are even considerably better than AgaA as reflected by the \( V_{\text{max}}/K_m \) values in Table 5 for hydrolysis at 37°C.

To verify that the described single-nucleotide replacements at position 355 are determinative for the new enzyme phenotypes, the observed point mutations were introduced into the \( agaB \) gene. This was carried out using our routinely applied \( E. coli \) expression vector construct (\( agaB \) in pBTac1) and a conventional site-directed mutagenesis (25). The resulting mutant enzymes were produced, isolated from \( E. coli \), and characterized. The mutants derived thereby were identical to AgaM4 and AgaM5, respectively (data not shown). The results indicate that amino acid residue 355 influences the kinetic properties and stability of the \( B. stearothermophilus \) \( \alpha \)-galactosidases. It seems unlikely that the modest increase in thermostability displayed by the mutants (Fig. 4B) is responsible for the dramatic difference in \( K_m \) values. It seems more likely that the residue at position 355 affects the affinity for the substrate more directly. A simple, plausible explanation may be that the binding or access to the binding site is obstructed by the large side chain of the glutamate residue but not by the smaller side chains of the residues found in AgaA (Ala) and the mutants (Gly and Val). Yet in absence of a three-dimensional structure of the enzyme, this remains speculative. According to the nature of the amino acid residue 355 in AgaA as well as in the thermoadapted mutants, nonpolar side chains or uncharged polar side chains of amino acid 355 may be compulsory for the enzyme to acquire the properties suitable for \( \alpha \)-galactosidase hydrolysis at high temperatures (60 to 70°C). Figure 5 shows an amino acid alignment of bacterial \( \alpha \)-galactosidase partial sequences which correspond to the \( BsmI-HindIII \) fragment of \( agaA \) and \( agaB \). All of them contain Ala in position 355 (according to the AgaB1 sequence numbering), except AgaB1 and the \( \alpha \)-galactosidase from \( Bacillus halodurans \) (accession number NP_243089), which contains Gly as observed in AgaM1 to AgaM4. Further analysis of the evolutionary potential of \( \alpha \)-ga-

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**TABLE 4.** \( K_m \) values for pNP-\( \alpha \)-galactosidase hydrolysis*

<table>
<thead>
<tr>
<th>Temp (°C)</th>
<th>( K_m ) (mM) for:</th>
<th>AgaA</th>
<th>AgaB1</th>
<th>AgaM4</th>
<th>AgaM5</th>
</tr>
</thead>
<tbody>
<tr>
<td>25</td>
<td></td>
<td>0.38</td>
<td>0.83</td>
<td>0.24</td>
<td>0.16</td>
</tr>
<tr>
<td>50</td>
<td></td>
<td>0.39</td>
<td>3.60</td>
<td>0.34</td>
<td>0.25</td>
</tr>
<tr>
<td>60</td>
<td></td>
<td>0.40</td>
<td>9.5</td>
<td>0.43</td>
<td>0.26</td>
</tr>
</tbody>
</table>

*Activity tests were done in triplicate. The maximum variation from the mean values (shown) was less than 5%.

**TABLE 5.** Kinetic parameters of \( \alpha \)-galactosidases

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>AS 355</th>
<th>Melibiose</th>
<th>Raffinose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( K_m ) (mM)</td>
<td>( V_{\text{max}} ) (U mg(^{-1}) mg)</td>
<td>( V_{\text{max}}/K_m )</td>
</tr>
<tr>
<td>AgaA</td>
<td>Ala</td>
<td>3.3</td>
<td>45</td>
</tr>
<tr>
<td></td>
<td>Glu</td>
<td>13.3</td>
<td>370</td>
</tr>
<tr>
<td>AgaB1</td>
<td>Gly</td>
<td>4.1</td>
<td>255</td>
</tr>
<tr>
<td></td>
<td>Val</td>
<td>3.2</td>
<td>83</td>
</tr>
</tbody>
</table>

*Hydrolyzing reactions were performed at 37°C in 0.1 M potassium phosphate buffer (pH 6.5). Michaelis-Menten kinetics of hydrolyzing reactions were verified by plotting reaction rates against substrate concentration. The kinetic parameters were determined graphically from Lineweaver-Burk plotting of the initial cleavage rate. Activity tests were done in triplicate. The maximum variation from the mean values (shown) was less than 5%. AS 355, amino acid residue at position 355.
lactosidases may be attempted, e.g., by applying this system for selection of mutants at temperatures over 70°C.

Concluding remarks and general discussion. Thermophilic selection systems, such as the one described in this paper, can be useful for isolation of enzyme variants with improved thermostability. Yet, substantial thermostabilization of mesophilic proteins by means of in vitro or in vivo evolution systems may be a difficult task since the routes around multidimensional thermodynamic obstacles to suitable thermostable structures are difficult to find and evolving structures tend to get caught in thermodynamic troughs. Another potential application for thermophilic in vivo evolution systems is to establish thermostable enzyme variants with improved specific activity at elevated temperatures for relevant substrates. In fact, this may be especially pragmatic since thermophilic enzymes sometimes show lower specific activity than their mesophilic or less thermophilic counterparts at their respective temperature optima. We have seen this in the case of α-galactosidases where the thermostable α-galactosidases from *Thermus* spp. display, in general, a lower specific activity for some substrates than their less thermostable counterparts from *B. stearothermophilus*, *E. coli*, and *Streptococcus mutans* at respective temperature optima (7; also unpublished results). Yet do we need a thermophile to promote such mutants; i.e., can an *E. coli* selection system serve this purpose? We have observed that selection of enzyme variants with improved catalytic activity can be dependent on the growth temperature of the selector organism; e.g., in previous work in this laboratory, an attempt was made to isolate variants of AgaB with improved catalytic activity by applying mutagenesis, using *E. coli* and selecting for growth at 37°C. Enzyme variants were isolated which displayed improved catalytic activity by site-directed mutagenesis are restricted. In this study we employed an in vivo system to investigate the evolutionary potential of an α-galactosidase. We succeeded in isolating mutants which supported the growth of the thermophilic host cells at an elevated temperature (67°C) on a selective medium. Characterization of the mutants revealed improved substrate affinity and catalytic capacity at elevated temperatures besides slightly enhanced thermostability. The results show that thermophilic bacteria can principally be applied for a thermoadaptation of glycoside-hydrolyzing enzymes. Thus, such in vivo evolution systems should be applicable to other industrially important enzymes as well, and in combination with in vitro evolution techniques, such as gene shuffling, should present a powerful method for obtaining thermoactive and thermostable enzymes. The prerequisite is the capability of the thermophilic host to import and metabolize the relevant substrates and the evolutionary potential of the particular enzyme.

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REFERENCES


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SELECTION OF α-GALACTOSIDASE MUTANTS 3391
ERRATA

ModE-Dependent Molybdate Regulation of the Molybdenum Cofactor Operon moa in Escherichia coli

Lisa A. Anderson, Elizabeth McNairn, Torben Lubke, Richard N. Pau, and David H. Boxer

Department of Biochemistry, University of Dundee, Dundee DD1 5EH, Scotland, and Nitrogen Fixation Laboratory, John Innes Centre, Norwich NR4 7UH, United Kingdom


Thermoadaptation of α-Galactosidase AgaB1 in Thermus thermophilus

Olafur Fridjonsson, Hildegard Watzlawick, and Ralf Mattes

Institut für Industrielle Genetik, Universität Stuttgart, 70569 Stuttgart, Germany

Volume 184, no. 12, p. 3385–3391, 2002. Page 3390: Figure 5 should appear as shown below.

AgaB1: DREBPILYMEKLDDNEEKLVNIKTEABGICHELDGGWFSKDDDRSLGWMHINR
AgaA1: DREBPILYMEKLDDNEEKLVNIKTEABGICHELDGGWFSKDDDRSLGWMHINR
AgaN B.stearotherm: DREBPILYMEKLDDNEEKLVNIKTEABGICHELDGGWFSKDDDRSLGWMHINR
RafA E.coli: QKLFYHIAWEGFINHPDYIMQMBERAAALLGRVE1IDDDGFKNRFAALLGTYDE
Aga S.mutans: NKEFPILYMEKLDDNEEKLVNIKTEABGICHELDGGWFSKDDDRSLGWMHINR
MelA T.ethanolicus: DKEFPILYMEKLDDNEEKLVNIKTEABGICHELDGGWFSKDDDRSLGWMHINR
AgaR P.pentosaceus: NQERRYPVYMEKLDDNEEKLVNIKTEABGICHELDGGWFSKDDDRSLGWMHINR
α-Gal Y.pestis: EKREPVYMEKLDDNEEKLVNIKTEABGICHELDGGWFSKDDDRSLGWMHINR
α-Gal S.pneumoniae: KKEFPILYMEKLDDNEEKLVNIKTEABGICHELDGGWFSKDDDRSLGWMHINR
α-Gal B.halodurans: DREBPILYMEKLDDNEEKLVNIKTEABGICHELDGGWFSKDDDRSLGWMHINR