Advantages and disadvantages of His-tagged beta-galactosidase

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Abstract

β-Galactosidase is one of the most important biotechnological enzyme used in the dairy industry, pharmacology and in molecular biology. In our laboratory we have overexpressed a recombinant β-galactosidase in *Escherichia coli* (*E. coli*). This enzyme differs from its native version (β-GalWT) in that 6 histidine residues have been added to the carboxyl terminus in the primary sequence (β-GalHis), which allows its purification by immobilized metal affinity chromatography (IMAC). In this work we compared the functionality and structure of both proteins and evaluated their catalytic behavior on the kinetics of lactose hydrolysis. We observed a significant reduction in the enzymatic activity of β-GalHis with respect to β-GalWT. Although, both enzymes showed a similar catalytic profile as a function of temperature, β-GalHis presented a higher resistance to the thermal inactivation and evidenced greater half-life time compared to β-GalWT. At room temperature, β-GalHis showed a fluorescence spectrum compatible with a partially unstructured protein however, it exhibited a lower tendency to the thermal-induced unfolding with respect to β-GalWT. Analytical ultracentrifugation experiments demonstrated that the population of β-GalHis molecules exhibited a higher proportion of monomers and a lower proportion of tetrameric species with respect to the His-tag free protein. The impairment of tetramerization may would explain the negative effect of the presence of His-tag on the enzymatic activity. In addition, the present results, analyzed in the context of the available literature, suggest that the effect of the His-tag is protein-specific.

Keywords
beta-galactosidase, His-tag, enzymatic activity, protein structure, thermal stability.

1. Introduction
β-D-galactosidase [EC 3.2.1.23] (β-Gal) is a glycoside hydrolase enzyme that catalyzes the hydrolysis of glycosidic bonds, producing monosaccharides from β-galactosides. β-Gal has been extensively studied because of its nutritional, biotechnological and therapeutic impact. In our laboratory, we are interested in describing how the recombinant production of proteins, particularly β-Gal, and the purification strategies could modulate their structure/function relationship.

Within the vast alternatives of biological systems for recombinant protein production, *E. coli* is the preferred microorganism for protein expression because of the easy handling and storage. Besides, in the protein purification step, histidine tags (His-tag) have gained great popularity. His-tags (typically containing six or more consecutive histidine residues) may be used for protein purification by immobilized metal affinity chromatography (IMAC), where divalent cations (usually Ni\(^{2+}\) or Co\(^{2+}\)) are adsorbed in an agarose matrix. The first report on using the nickel coupled to nitriloacetic acid agarose system (Ni\(^{2+}\)-NTA) for His-tag protein purification was dated on 1975. His-tag protein goes through the agarose column and attach to it. In principle, the basic nitrogen atom in the imidazole of histidine has an alone electron pair that coordinate with metallic cations (Ni\(^{2+}\)). Also, nonspecific interactions, e.g., electrostatic and hydrophobic forces contribute to protein-agarose binding. Then histidine or imidazole have been applied as elution agents for immobilized metal affinity displacement chromatography.

In some cases, the tags also help in protein expression, folding, and/or solubility. For instance, *in vitro* designed oligomeric proteins have been engineered through the interaction of His-tag with divalent cations as protein biomaterial with novel therapeutic properties. Moreover, a family of metal-NTA based fluorescence probes has been developed for the intracellular visualization of His-tag proteins. Immobilized metal affinity (IMA) strategy was also used to study protein-membrane interaction. Raghunath et al. studied the kinetics of His-tag protein binding to nickel decorated liposomes. They found that protein-membrane association involves the binding and reorganization of protein conformation according to an increase in the membrane packing. This approach allows to describe the intermediates preceding membrane bending driven by protein crowding.
It is usually accepted that due to their small size, His-tags would not interfere with the function and structure of the majority of proteins \(^{10,15}\). However, there is an increase number of reports showing that this conception may be not always true \(^{16-18}\). Moreover, there are evidences that His-tags may affect the oligomeric states of proteins as well as their function \(^{19-21}\).

In the present study we compared a β-Gal containing a hexahistidine peptide added in the C-terminal, β-Gal\(_{\text{His}}\), with its wild type commercial counterpart, β-Gal\(_{\text{WT}}\). Our results contributed to understand the unique effects of His-tag in proteins.

2. Materials and Methods

2.1. Materials.

Luria-Bertani medium (LB). Isopropyl-β-D-thiogalactopyranoside (IPTG), Kanamycin Sulfate (Sigma Aldrich); *Escherichia coli* K-12 W3110, *Escherichia coli* strain BL 21\(\lambda\)Codon plus (Novagen).

Other reagents and solvents were of analytical grade.

2.2. Methods.

2.2.1. Cloning β-Gal gene.

The β-galactosidase gene was amplified from *E. coli* K-12 W3110 genomic DNA by PCR using primers Bgal S1N (5’-CATATGACCATGATTACGGATCTGGCCGTGCATTACAAAACGTCG-3’) and BgalA1 (5’- GGCTCGAGTTTTGACACCAGAACCAGACCTGGTA-3’) the PCR products were partially digested with NdeI and Xho I and inserted into pET26 (+) using the same restriction enzymes, yielding the plasmid pET 26+βGal which contains a C-terminal six histidine tag sequence\(^{22}\).

2.2.2. β-Gal Expression and Purification.

*Escherichia coli* strain BL21\(\lambda\)Codon plus (Novagen) transformed with Plasmid pET 26+βgal was grown at 37°C in Luria-Bertani medium (LB) supplemented with kanamycin 50 µg/ml\(^{22}\). β-Gal expression was induced adding isopropyl-β-D-thiogalactopyranoside (IPTG) to a final concentration of 0.4 mM at OD\(_{600}\) of approximately 0.6 and further incubation at 20°C for 20 h. The grown cells were harvested by centrifugation at 5,000 g for 20 min at 4°C, resuspended in 20 mL of NaCl 500 mM in phosphate buffer 0.05 M, pH 6.8. Then the cells were incubated with lysozyme (0.3 mg/mL) for 30 min. at 20°C. After that, bacteria were disrupted by three cycles of sonication of 30 seconds each, frozen with liquid nitrogen.
and thawed in a water bath at 35°C. Then, the lysed cells were centrifuged at 10000 rpm for 30 min. at 4°C and the supernatant was then incubated with 1 mL Ni-NTA beads (ProBond™ resin Invitrogen) for 3h at 4°C by tumbling. Then the beads were poured in cartridges and washed with imidazole solution of 0, 60, 100, 200 and 400 mM. Each elution sample was dialyzed against 0.05 M phosphate buffer (pH 6.8) at 4°C for 20 h with 3-4 changes of buffer.

The sample eluted with the 200 mM imidazole contained a considerable amount of highly pure β-Gal tested by SDS-PAGE and finally the pure enzyme was stored at 4°C in dialysis buffer (at 3 mg/mL).

β-Gal SDS-PAGE

The β-Gal purity was evaluated by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) (10%). Samples were heated at 100 °C during 5 min. in loading buffer. The amount of protein submitted to SDS-PAGE was around 90 µg. After running, the proteins in the gel were stained with coomassie blue.

2.2.4. Protein quantitation.

Protein concentration was measured by the Lowry’s method.

2.2.3. β-Gal activity assay.

2.2.3.1. Determination of kinetic parameters.

The reaction was carried out in phosphate buffer 0.05 M, pH 6.8 using monohydrated Lactose (Anedra) as substrate. It was initiated by the addition of 0.01 mL of the enzyme preparation containing (0.001 mg to 0.01 mg of β-Gal) and incubated for 20 min at 37°C. Then, the reaction was quenched by boiling 5 min. β-Gal activity was measured by quantitative analysis of the glucose released, determined by the glucose oxidase method. One unit of enzyme activity was defined as the amount of enzyme required to produce 1 µmol of glucose/min in the described conditions.

The values of $K_M$ and $V_{max}$ were determined by fitting the Michaelis–Menten equation to the $V_0$ versus substrate concentration data plot by a computer aided nonlinear regression analysis.
2.2.3.2. Temperature-dependent β-Gals activity profile.

The thermal activity profiles of β-Gal\textsubscript{WT} and β-Gal\textsubscript{His} were evaluated from 25°C to 65°C at pH 6.8. The other reaction conditions were described above.

2.2.3.3. Thermal stability of β-Gals.

The thermal stability of β-Gal\textsubscript{WT} and β-Gal\textsubscript{His} was studied by measuring the residual catalytic activity determined as described above after a pre-incubation period of 20 min. at different temperatures (20 - 60°C).

2.2.3.4. Half-life time.

Residual activity was measured as described above, a closed cap tube containing 3 µl of β-Gal (0.01 g/L), preincubated at 50°C during a time ranging from 2 to 20 min. After this time, 300 µl of lactose (200 mM) was added and the specific enzyme activity was measured as described above at 37°C. The profile of residual activity was plotted vs time.

A single exponential decay mathematical approach was applied to fit the curve of specific activity (SA) vs time, in order to determine the half-life time ($\tau_{1/2}$).

\[
SA_{0.5} = SA_{\text{initial}} \times e^{-k} \quad (1)
\]

\[
\frac{SA_{0.5}}{SA_{\text{initial}}} = e^{-k \tau} \quad (2)
\]

\[
\ln 0.5 = -k \tau_{1/2} \quad (3)
\]

\[
\tau_{1/2} = \frac{0.69}{k} \quad (4)
\]

where \(SA_{\text{initial}}\) is the specific activity measured at zero time, \(SA_{0.5}\) is the half value of \(SA_{\text{initial}}\), \(k\) is the constant of exponential decay, and \(\tau_{1/2}\) is the half-life time.
2.2.5. Structural properties of β-Gals.

2.2.5.1. Fluorescence spectroscopy.

Fluorescence spectra were recorded in a Fluoromax Spex-3 JovinYvon (Horiba, New Jersey, USA) spectrofluorimeter. A quartz cell with 10 mm path length and a thermostated holder was used. The slits and λ_ex were set at 2 nm and 290 nm, respectively. Emission spectra were acquired within the 300-400 nm range. Protein concentration used was 0.2 mg/ml. Raman scattering contribution from water was subtracted in all spectra. To facilitate comparisons, λ_max was determined but also the center of spectral mass (CSM) was calculated for each fluorescence emission spectra \(^{26}\) according to equation (5), where \(I_i\) is the absorbance or the fluorescence intensity measure at the wavelength \(\lambda_i\).

\[
\lambda = \frac{\sum \lambda_i I_i}{\sum I_i} \tag{5}
\]

We also analyzed the λ_max from each protein vs temperatures within 20 to 60 °C range. At rate heating of 1°C/min.

2.2.5.2. Analytical ultracentrifugation (AUC).

β-Gals (0.5 mg/mL) solutions prepared in 0.05 M phosphate buffer, pH 6.8 were submitted to Sedimentation Velocity (SV) analysis as described previously \(^{27}\). Briefly, it was carried out with an Optima XL-A analytical ultracentrifuge (Beckman Coulter Inc., Fullerton, CA, USA) equipped with a spectrometric UV-visible system. All SV experiments were conducted at 40,000 rpm at 20°C; 50 scans were recorded along 2 h, at 280 nm. The reference cell contained buffer solution. To obtain the sedimentation coefficient (S) of each species, data were analyzed with SEDFIT software, using the continuous sedimentation coefficient distributions c(s) model \(^{28}\). The experimental s-value were transformed into a value under standard conditions (water 20°C). The viscosity and density values (required for S calculations) in the case of β-Gal solutions were determined using SEDNTERP software \(^{29}\). S has the dimensions of time units expressed in Svedberg (1S=10^{-13}s) and can be calculated by equations (6) and (7).
\[ v_t = \frac{M \cdot r \cdot \omega^2}{6 \pi \cdot r_0} \]  

(6)

\[ S = \frac{v_t}{r \cdot \omega^2} = \frac{M}{6 \pi \cdot r_0} \cdot v_t = \frac{m \cdot r \cdot \omega^2}{6 \pi \cdot r_0} \]  

(7)

where \( v_t \) is the sedimentation rate, \( r \) is the distance of the particle from the axis of rotation, \( M \) the molar mass of the particle and \( \omega \) is the angular velocity of the rotor equation (6). \( S \) serves to normalize the \( v_t \) of a particle by the acceleration applied to it \( (r \cdot \omega^2) \). The resulting value is independent on the acceleration but depends on the properties of the particle (the mass \( m \) and the hydrodynamic radius \( r_0 \)) and the viscosity of the medium (\( \eta \)) where it is suspended as shown by equation (7). Experimentally, the absorbance profile obtained by analytical ultracentrifugation is described by the Lamm equation equation (8) and \( S \) can be calculated:

\[ \frac{d[\beta Gal]}{dt} = \frac{1}{r} \frac{d}{dr} \left[ rD(M) \frac{d[\beta Gal]}{dr} - S(M) \omega^2 r^2 [\beta Gal] \right] \]  

(8)

where \( D(M) \) is the diffusion coefficient which, as well as \( S(M) \) is dependent on the molar mass \( (M) \) of the particle (in this case the \( \beta \)-Gal molecule).

2.6 Statistical calculations.

The least squares method was applied to fit functions through nonlinear regression analysis. Pairwise comparisons were made with the Student’s t-test using Sigma Plot version 12.5.

3. Results and Discussion.

3.1. \( \beta \)-Gals functionality.

In most cases, the incorporation of His-tag has resulted in great increases in the efficiency of the protein purification process however, sometimes, it hampers the protein functionality. Then, several experiments were conducted to remove the additional histidine residues but drawbacks remain if non-wild type amino acids remain in the protein\(^{18}\). In our experiments we evaluate the enzymatic activity of the wild type \( \beta \)-Gal\(_{WT}\), the commercial enzyme from Sigma, and a recombinant \( \beta \)-Gal\(_{His}\) that we
produced and purified by IMAC chromatography. Both enzymes show a Michaelian behavior. However, 
β-GalHis exhibits lower specific activity with respect to β-GalWT (Fig.1, Table 1); a $V_{\text{max}}$ decrease of 
around 60% is observed for β-GalHis with respect to β-GalWT. On the other hand, the presence of histidine 
residues also seems to moderately favor the affinity of the active sites for lactose ($\beta$-GalHis$K_M < \beta$- 
GalWT$K_M$), although, the difference observed in the later parameter was not statistically significant. We 
ruled out the possibility that the difference in the kinetic parameters were due to impurities in the protein 
sample. SDS-PAGE (Fig 1.b) showed that both samples only present the β-gal monomer (116 kD) band. 

![Functional β-Gal properties](image)

**Fig. 1 Functional β-Gal properties.** a) Specific activity as a function of lactose concentration ($\beta$-GalWT 
white symbols, β-GalHis, black symbols). Samples were incubated for 20 min, at 37°C and pH 6.8 and 
all measurements were carried out in conditions of initial velocity, according to Michaelis-Menten 
(MM) model. Hyperbolic curves could be adjusted to the experimental points, according to the MM 
equation. The resulting kinetic parameters are shown in Table 1.b) β-GalS SDS-PAGE.

**Table 1. Kinetic and structural parameters for β-GalHis and β-GalWT**

|                      | β-GalHis | β-GalWT |
|----------------------|----------|---------|
| **Kinetic parameters** |          |         |
| $V_{\text{max}}$ (µmol/min/mg) | 0.163 ± 0.005* | 0.46 ± 0.02* |
| $K_M$ (mM)            | 7.27 ± 0.73 | 9.9 ± 1.5 |
| **Structural parameters** |          |         |
| $\tau_{1/2}$ (min)    | 8.1 ± 1.2# | 5.6 ± 0.9# |
| $\lambda_{\text{max}}$ (nm) | 347       | 345     |
CSM (nm) | 360 | 356
---|---|---

Vmax, Km and \( \tau_{1/2} \) values are the mean ± sem. *statistically significant difference (P = <0.001). #statistically significant difference (P = 0.045).

Firstly, we decided to add the His-tag at the carboxyl terminus of β-Gal because the amino terminus of the protein takes part of the active site 30, in spite of the fact that Brome and her colleagues expressed an active β-Gal with the His-tag in the amino terminal of the primary sequence 31. Besides, Ullman 32 found β-Gal activity in hybrid proteins where tags appeared localized in both extremes of the sequence. In those cases, any structure-function analysis nor any comparison to the wild type protein was developed to describe the tags effects on the protein.

For other proteins, there are differences on the molecular performance according to the location of the His-tag within the protein sequence 33,34. For instance, the rat corticotropin-releasing factor receptor type 2a adopts two different disulfide bond patterns at the N-terminus when the His-tag is at either the N- or C-terminal 35. Otherwise, the effect of His-tag on protein function is varied according with the protein. In GNAT superfamily, the His-tag binds at the substrate site and acts as a weak competitive inhibitor for the substrate 17. On the other hand, the lower catalytic efficiency of the His-tagged with respect to the wild type chondroitinase ABC-I was related with the difficult to completely degrade the chondroitin sulfate to low molecular weight products 36.

Beyond the diminution we observed in the enzymatic activity, we investigated if the His-tag affected the thermal properties of β-Gal. For that, different approaches were used: a) the temperature effect on enzyme activity, b) the resistance to thermal inactivation and c) the half-life time for the enzyme denaturation. Results are shown in Fig. 2 and Table 1. Both enzymes exhibit the same catalytic profile vs reaction temperature with an optimal activity at around 45°C (Fig. 2a). Moreover, the negative effect of high temperature was much more important for the wild type than for the His-tagged enzyme: when the enzymatic activity is measured at 50°C, β-Gal_{His} maintains approximately 50% while β-Gal_{WT} only exhibits around 20 % of the optimal activity. In terms of thermal stability, β-Gal_{His} presents a higher
resistance to temperature inactivation compared to $\beta$-Gal$_{WT}$ as shown in Fig. 2b. In order to evaluate this difference, the half-life time of each enzyme (Fig. 2c) was studied. The preincubation of the enzymes at 50°C $\beta$-Gal$_{His}$ reflects this tendency and $\beta$-Gal$_{His}$ seems to be more stable with respect to $\beta$-Gal$_{WT}$ (Table 1). The effect of His-tag on proteins thermal stability is different depending on the protein but in general, a negative effect is observed. Chen and his collaborators 36 found that the presence of a His-tag produced a particular result in the choindroitinase ABC1 (ChSase ABC I) thermal response. Their results demonstrated that His-tag could improve the thermostability of ChSase ABC I when it was incubated at relative low temperatures (30°C-35°C), but the opposite effect occurs at high temperatures (40°C-45°C).

On the other hand, Booth and coworkers 18 using differential scanning fluorimetry studied the thermostability for a set of ten different N-Terminal His-tagged proteins and found that in almost all cases, the His-tagged version was less stable than the wild type protein.

This scenario encourages us to explore the structural features underlying the differential behavior of $\beta$-Gal$_{WT}$ and $\beta$-Gal$_{His}$. 
Fig. 2. Thermal and temporal inactivation profiles of β-Gals. a) Specific activity measured at the indicated temperature within the range 22°C - 65°C. b) β-Gal specific activity measured at 37°C after preincubation for 20 min. at different temperatures (20-60°C). In each data set normalization was done with respect to the corresponding specific activity preincubated at 30°C, which was taken as the unity. c) Catalytic activity was evaluated at 37°C, pH 6.8 and lactose 200 mM after preheating the proteins (0.01 g/L) at 50°C during a fix time period (20-60 min). Exponential decay curves (dashed lines) are the fitness to the experimental points (black and white circles) and allowed estimating the half-life times (shown in Table 1).
3.2. Structural properties of β-Gals.

3.2.1. Fluorescence Spectroscopy.

The fluorescence emission of Trp is sensitive to the polarity of the environment and can be used to detect conformational changes in proteins. β-Gal is a tetrameric protein with 39 Trps residues per protomer. β-GalWT fluorescence spectrum shows a λ\textsubscript{max} at 345 nm and CSM=356 nm (Fig. 3a, Table 2) which is compatible with Trp residues localized in highly polar environment if compared with the spectrum of Trp localized in a buried or non-polar medium (λ\textsubscript{max}~325 nm). It is noteworthy that we tested the Mg\textsuperscript{2+} free protein. This condition could contribute to the higher value of λ\textsubscript{max} in β-Gal\textsubscript{WT} spectrum compared to those previously reported for the same protein. For β-Gal\textsubscript{His}, the spectrum shows an important bathochromic shift, with a λ\textsubscript{max} at 347 nm and CSM=360 nm (Fig. 3a, Table 1). These results indicate that Trp residues became, on average, more accessible to the solvent and that the His-tag could induce at least a partial unfolding of β-Gal or, on average, a highly hydrated structure.

However, when λ\textsubscript{max} was evaluated along heating (Fig. 3b) we observe that the His-tagged protein shows a greater thermal stability with respect to the wild type protein (compare the magnitude of the thermal λ\textsubscript{max} increase up to 50°C in both samples). In our laboratory we have also described an increase in the thermal stability of β-Gal\textsubscript{WT} when interacting with model membranes or in molecular crowded conditions. The present results indicate that the covalent modification on β-Gal\textsubscript{WT} leading to β-Gal\textsubscript{His} also accounts for the increase resistance to inactivation that displays the latter compared with the former (Fig. 2b). Moreover, at 50°C and above we observe the major differences in the λ\textsubscript{max} value between both proteins. It is noteworthy that at 50°C β-Gal\textsubscript{His} shows lower values of λ\textsubscript{max} with respect to β-Gal\textsubscript{WT} and as consequence, an opposite structural behavior respect to what was observed at 25°C.
Fig. 3. Structural analysis of β-Gals a) Intrinsic fluorescence spectra of β-GalHis (dotted line) and β-GalWT (full line) proteins at 0.2 g/L at 25°C. b) Effect of temperature on the λ_{max} of β-GalHis (●) and β-GalWT (○).

The most active form of β-GalWT have been described when the protein oligomerized as a tetramer. It is well documented that the decrease of the tetramer in favor of minor oligomeric states could lead to a negative effect in β-Gal functionality. However, a superactive molecule could be observed when higher oligomeric and/or particulated forms of the enzyme are present in the sample. On the other hand, Wu and Filutowics proposed that His tagged proteins may differ from their wild type counterparts in dimerization/oligomerization and afterwards, many researchers lead to similar conclusion. Also, His-tag promoted the dimerization of HSC70 C30ΔL-His but had no effect on the elution profile of HSC70 C30WT-His(+), compared to their respective untagged forms. Moreover, Kenig and her colleagues demonstrated that the His-tagged proteins were located intracellularly as soluble proteins and also in an aggregated form as inclusion bodies. In contrast, the non-tagged proteins were found only in the soluble form and this fraction was used for further purification studies. With those concepts in mind we decided to evaluate the oligomeric state of our β-Gals.
3.2.2. Analytical ultracentrifugation (AUC)

In order to evaluate the effect of His-tag on the supramolecular organization of the enzyme we developed AUC. Both enzymes presented a heterogeneous profile, but the presence of histidine seems to increase the population of monomers (Table 2, Fig. 4). Besides, it was remarkable the lower proportion of the tetramers’ population found in Gal\textsubscript{His} if compared with \(\beta\)-Gal\textsubscript{WT}. These phenomena contribute to explain the negative effect of the presence of a His-tag on the enzyme activity.

![Analytic ultracentrifugation](image)

**Fig. 4. Analytic ultracentrifugation** of \(\beta\)-Gal\textsubscript{His} (full line) and \(\beta\)-Gal\textsubscript{WT} (dotted line)

**Table 2. Data from AUC hprofile from each \(\beta\)-Gal**

|                  | \(\beta\)-Gal\textsubscript{His} | \(\beta\)-Gal\textsubscript{WT} |
|------------------|----------------------------------|----------------------------------|
| **Supramolecular organization** | % | Size (kD) | % | Size (kD) |
| monomer          | 15.1 | 115.7     | 7.2 | 106.6     |
| tetramer         | 39.2 | 333.4     | **49.3** | 387.8     |
| Oligomer 1       | **20.7** | 626.7     | 16.7 | 656.9     |
| Oligomer 2       | 8.6  | 998.9     | 8.5  | 904.3     |
| Oligomer 3*      | 16.1 | 1300-3001 | 18.2 | 1141-1740 |

*Correspond to the sum of all the oligomers bigger than 1141 kD that appear in the AUC data
Another important finding was the lower size (or hydrodynamic radius) of the β-Gal_{His} tetramer if compared with that of β-Gal_{WT} which may also be associated with an inactive conformational structure of the protein. All these results could explain the decreased in His-tagged protein functionality. This statement is supported by the fact that the active site is made up of elements from two subunits of the tetramer, and disassociation of the tetramer into dimers removes critical elements of the active site^{30}.

4. Conclusions

In the present work, through kinetic, fluorometric and ultracentrifugation analysis, we demonstrated that the addition of a His-tag to E. coli β-Gal reduces the catalytic activity possibly through an impairment of the proper acquisition of the typically active quaternary structure (tetramers). However, the tendency to form higher oligomeric structures may explain the improvement in the thermal structural and functional stability. It is important to consider that, in view of the present study and the available literature, the effect of His-tag seems to be protein-specific.

Declarations

Ethics approval and consent to participate

Not applicable

Consent for publication

Not applicable

Availability of data and materials

The datasets during and/or analysed during the current study available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests

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