Isolation and characterization of the mutant form of N-terminal catalytical module of Bos taurus tyrosyl-tRNA synthetase with the replacement of Trp 40 and Trp 283 by alanine

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Aim. To obtain the mutant one-tryptophan form of the N-terminal catalytical module of Bos taurus tyrosyl-tRNA synthetase (mini TyrRS) for the monitoring of the local conformational changes by fluorescence spectroscopy. Methods. Bacterial expression, metal-chelating chromatography, gel electrophoresis, fluorescence spectroscopy, computational modeling. Results. The replacement of two tryptophan codons with alanine codons in cDNA encoding mini BtTyrRS has been performed. These mutations do not affect the synthesis and solubility of the mini BtTyrRS in E. coli BL21 (DE3) pLysE strain. The amount of a soluble form of the mutant mini BtTyrRS in the cytoplasm of bacterial cells at the incubation of bacterial culture at 25 °C was about 47 % of the total amount of recombinant protein. Computational modeling and fluorescence study of the single-tryptophan form of mini BtTyrRS revealed that Trp 87 residue was localized at the dimerization region of the enzyme. The characteristics of tryptophan fluorescence of the mutant mini BtTyrRS indicated that Trp 87 is localized in the immobilized microenvironment of the dimer interface. Conclusions. The optimal conditions of bacterial expression of the mutant Trp 87-containing form of mini BtTyrRS in the bacterial culture of E. coli BL21 (DE3) pLysE strain have been established. The Trp 87-containing form of mini BtTyrRS is suitable for monitoring the local conformational changes at the enzyme dimer interface.

Keywords: tyrosyl-tRNA synthetase, mini TyrRS, bacterial expression, fluorescence spectroscopy, computational modeling
Introduction

Aminoacyl-tRNA synthetases (ARSases) [EC 6.1.1.1] are the key enzymes of protein biosynthesis. At the preribosomal stage of translation, ARSases catalyze the activation and covalent attachment of the amino acids to the homologous transfer RNAs, thus carrying out the initial stage of the realization of genetic information into the protein structure [1, 2].

Mammalian tyrosyl-tRNA synthetase (TyrRS, EC 6.1.1.1) is one of the most studied eukaryotic ARSases. Under physiological conditions, *Bos taurus* tyrosyl-tRNA synthetase is an α2 homodimer with a molecular weight of 2×59.2 kDa. Each monomer consists of two structural parts: the N-terminal catalytic form (mini *Bt*TyrRS, 39 kDa) and the C-terminal EMAP II-like module (20 kDa) [2]. During the isolation of TyrRS from bovine liver, it was shown that along with the full-length main form, the functional proteolytically modified form of tyrosyl-tRNA synthetase with a molecular weight of 39 kDa was released, which retains its enzymatic activity *in vitro* [3, 4].

In addition to the basic tRNA aminoacylation function the mammalian tyrosyl-tRNA synthetases perform also the non-canonical functions: after enzymatic cleavage of tyrosyl-tRNA synthetase by elastase into mini *Bt*TyrRS, which are respectively located in the active center of the enzyme (W40), in the region of dimerization of mini *Bt*TyrRS monomers (W87) and in the binding site of the tRNA anticodon triplet (W283). Such location of tryptophan residues in the functionally important regions of the protein’s amino acid sequence makes it very promising to study the properties by fluorescence spectroscopy, especially if there is only one residue in one of three positions in the enzyme structure. Previously, we have cloned the cDNA of the tyrosyl-tRNA synthetase catalytic module in the expression plasmid pET32a and investigated its expression [9]. Subsequently, Trp40 and Trp283 codons were replaced with alanine codons by site-directed mutagenesis in cloned cDNA and only one tryptophan codon was left at the site of dimerization of mini *Bt*TyrRS monomers [10].

The purpose of this work was to determine the optimal expression conditions of mutant mini *Bt*TyrRS and to isolate the recombinant protein for further study of its properties, especially the local conformational changes at the enzyme dimer interface.
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**Materials and Methods**

In order to obtain the mutant form of *Bos taurus* mini TyrRS we used the bacterial expression system of *E. coli* cells [11–13]. The synthesis of recombinant protein was carried out in the *E. coli* strain BL21 (DE3) pLysE (Stratagene, USA) transformed with pET30a-39KYSW87 plasmid.

The pET30a-39KYSW87 expression construction was created on the basis of the pET30a(+) vector (“Novagen”, USA) and contained a cloned cDNA with the tryptophan-40 and tryptophan-283 codons replaced by alanine codon. A plasmid DNA was isolated using the Gene JET Plasmid Miniprep Kit from “Thermo Scientific”. The concentration of plasmid DNA was determined on NanoDrop 2000 spectrophotometer (“Thermo Scientific”).

In order to obtain the recombinant plasmid construct pET30a-39KYSW87, transform it into *E. coli* cells and express the mutant cDNA of the catalytic module *B. taurus* tyrosyl-tRNA synthetase, the genetically engineered *E. coli* DH5α and BL21 (DE3) pLysE strains were used. Competent *E. coli* cells were obtained according to the Inoue method [14]. All procedures for transformation of plasmid pET30a-39KYSW87 into competent *E. coli* cells and analysis of plasmid by 0.7–1 % agarose gel electrophoresis were performed according to [15].

The cultivation of *E. coli* BL21 (DE3) pLysE cell culture and the induction of expression of recombinant mini *Bt*TyrRS in bacterial culture were performed in Luria-Bertani medium (LB) with 30 μg/ml kanamycin. Transformed with recombinant plasmid pET30a-39KYSW87, the competent *E. coli* BL21 (DE3) pLysE cells were grown on a shaker (BioSun Shaker Incubator ES-20) at 37°C to an optical density of $A_{600} = 0.6–0.8$ and the target protein synthesis was induced by adding 1M isopropyl-β-D-thiogalactopyranoside (IPTG) up to 1 mM concentration followed by incubation at 37°C for 4 hours and at 30°C and 25°C for 12 hours. The collected biomass from 100 ml of culture was resuspended in 12 ml of cell lysis buffer (50 mM sodium phosphate buffer, pH 8.0, 500 mM NaCl, 10 mM imidazole, 5 mM β-mercaptoethanol). Cells lysis was performed using an ultrasonic disintegrator (6 cycles of 20 s, 20 s intervals). The lysate was clarified by centrifugation at 13000 rpm for 30 min at 4°C.

The supernatant was applied to a Ni-NTA agarose column previously washed with 10 ml of washing buffer (50 mM sodium phosphate buffer, pH 8.0, 500 mM NaCl, 20 mM imidazole, 5 mM β-mercaptoethanol) and with lysis buffer. Recombinant protein was eluted with 5 ml of elution buffer (50 mM sodium phosphate buffer, pH 8.0, 150 mM NaCl, 200 mM imidazole, 5 mM β-mercaptoethanol). All protein containing fractions were combined and dialyzed against 500 ml of dialysis buffer (500 mM sodium phosphate buffer pH 8.0, 150 mM NaCl) for 20 hours at +4°C. The concentration of purified mini *Bt*TyrRS mutant protein was determined spectrophotometrically on BioMate-5 spectrophotometer using a molar extinction coefficient of 27850 M⁻¹ cm⁻¹ at a wavelength of 280 nm. The optical absorption coefficient of mini *Bt*TyrRS was determined by amino acid analysis of protein using ProtParam server (http://expasy.ch/cgi-bin/protparam). According to the ProtParam server, the molecular weight of the obtained recombinant
mutant mini $Br$TyrRS is 47364.36 Da and isoelectric point pI = 6.42.

Expression of the recombinant proteins was analyzed by SDS-PAGE [16].

Gels were stained with Coomassie blue R250 dye. The estimation of the amount of recombinant protein in the precipitate and in the supernatant fraction on the electrophogram was performed densitometrically on a ChemiDocTM XRS + system instrument (“BioRad”, USA).

The amino acid sequence of Bos taurus TyrRSs was used from the NCBI Gene database (https://www.ncbi.nlm.nih.gov/protein/) with identification numbers DAA32266.1. The three-dimensional crystal structures were obtained from RCSB PDB archive. Three-dimensional coordinates of the protein structural templates were obtained from Protein Data Bank (PDB) (http://www.pdb.org/pdb). Visualization and analysis of protein structure were performed using UCSF Chimera software [17]. The spatial structures of the $Br$TyrRS dimer and double mutant $Br$TyrRSW87 were modelled from the crystal structures of HsTyrRS (PDB codes 1N3L:A) as templates using SWISS-MODEL web-server [18]. High-resolution protein structure refinement was done by ModRefiner [19]. The final model of $Br$TyrRS structure was validated by the MolProbity server [20].

All fluorescence spectra were recorded at 25 °C on a Hitachi Model 850 fluorescence spectrophotometer equipped with thermostat-ed cell holder (Hitachi Ltd., Japan). Fluorescence measurements were performed in a quartz cell with an optical path length of 0.5 cm. The temperature in the quartz cuvette was determined within $\pm$ 0.2 °C. Both excitation and emission slits of 5 nm were used in all fluorescence measurements. The excitation light wavelength was 280 nm or 295 nm, the wavelength interval for the fluorescence spectra was 300–400 nm and the fluorescence registration was performed at the 90° angle to the beam direction.

Results and Discussion

Previously, we have cloned and sequenced the complete nucleotide sequence of cDNA of the Bos taurus tyrosyl-tRNA synthetase gene [2]. Based on cDNA, an expressive plasmid construct of pET-30a (+)-39KYRS was created with a cloned sequence of the synthetase N-terminal catalytic module. Expressed in strain E. coli BL21 (DE3) recombinant mini $Br$TyrRS retained the aminoacylating ability inherent in the native aminoacyl-tRNA synthetase.

Based on site-directed mutagenesis, the recombinant plasmid pET-30a (+)-39KYRS was used to create the substitutions of tryptophan residues at positions 40 and 283 by alanine in the cDNA of the synthetase catalytic module [10]. The resulting plasmid pET-30a-39KYRSW87 having only one tryptophan codon in the cloned cDNA was used in this work to obtain one-tryptophan protein for further fluorescence studies of conformational features and intramolecular interactions in protein structure [21, 22]. The amino acid alanine was selected for site-directed mutagenesis due to its small hydrophobic radical, which does not affect a secondary protein structure formed by the adjacent amino acid residues in the polypeptide chain.

A considerable amount of recombinant protein is required to investigate the properties of
the enzyme by experimental methods. Since the final yield of the target recombinant protein in bacterial systems strongly depends on the culture conditions, the experimentally determined optimal parameters for the expression of mini BrTyrRS in E. coli are required [9,10].

The native mini BrTyrRS cloned in plasmid pET-30a-39KYRS was used to determine the conditions of its optimal expression in the E. coli system [9]. It was shown that the highest level of synthesis of recombinant mini BrTyrRS in E.coli culture was achieved by adding to the culture medium of IPTG at a concentration of 1mM in the logarithmic phase of growth of the culture when it reaches an optical density OD600 = 0.7–0.9 for the induction of protein synthesis and incubation of the culture for 4 hours at 37° C. It was found that the composition of the culture medium had no significant effect on the expression of the enzyme.

We used these experimentally established optimal expression conditions to express and obtain a mutant single-tryptophan form of mini BrTyrRS in transformed plasmid pET30a-39KYRSW87 E. coli cell culture of strain BL21 (DE3) pLysE.

It is known that the sequence of the cloned genes in the expression vectors of the pET series plays a significant role in both the synthesis of recombinant proteins and obtaining soluble fraction of newly synthesized proteins [13]. Therefore, the replacement of two tryptophan residues with alanine in the mini BrTyrRS structure in our case could be critical and lead to a decrease in enzyme synthesis or its transition into insoluble inclusion bodies. In this regard, we simultaneously expressed in E. coli mutant and native mini BrTyrRS forms of plasmids pET-30a (+) — 39KYRS and pET30a-39KYRSW87. Our preliminary electrophoretic data showed that the mutations did not affect the expression of the mutant protein: the number of native and mutant forms of the tyrosyl-tRNA synthetase catalytic module synthesized in bacterial cultures was almost the same. However, our analysis of cell precipitate after clarification of bacterial lysates in the process of protein isolation showed that the majority of both native and mutant forms of the enzyme are in the cytoplasm in insoluble fraction of the inclusion bodies (results not shown).

The temperature of incubation is one of the major factors affecting the transition of recombinant proteins during expression in E. coli to the aggregated state, especially in vectors of the pET series with the extremely strong RNA polymerase promoter of phage T7 [13, 23, 24]. Therefore, it was decided to analyze the expression of the target protein in E. coli culture of strain BL21 (DE3) pLysE after IPTG induction at a lower incubation temperature of 30 °C and 25 °C. The results of the analysis are shown in Fig. 1 (A, B). The experimental data showed that with decreasing temperature of the bacterial culture growth, the amount of synthesized recombinant protein in the soluble fraction increased in proportion to the temperature decrease. The highest amount of recombinant mini BrTyrRS in soluble cytoplasmic cell fraction was obtained at the incubation temperature of 25 °C. At this temperature, the soluble fraction was about 47 % of the total amount of recombinant protein synthesized, whereas at 37° C it was only 13 %.

The established conditions for optimal expression of the catalytic N-terminal module of the Bos taurus tyrosyl-tRNA synthetase in
*E. coli* in LB medium were taken into account when obtaining a preparative amount of recombinant mutant mini *Br*TyRS in *E. coli* strain BL21 (DE3) pLysE using metal chelating chromatography. After lysis of bacterial cells by sonication and chromatographic purification of lysate on Ni-NTA agarose from 100 ml of the bacterial culture incubated at 25 °C for 8 hours we could get up to 3 mg of homogeneous recombinant protein of the mutant mini *Br*TyRS, with the purity according to electrophoresis about 95 % (Fig. 2). Up to 30 mg of purified recombinant enzyme can be obtained from 1 liter of bacterial culture under certain conditions of expression. Taking into account the aggregated protein in the inclusion bodies, the total yield of the synthesized recombinant mini *Br*TyRS in transformed plasmid pET30a-39KYRSW87 strain *E. coli* BL21 (DE3) pLysE was about 75 mg from the 1L LB medium.

According to the ProtParam program analysis, both native and mutant mini *Br*TyRS proteins are stable structures. Their instability indexes are almost identical, 36.2 and 37.15, respectively, indicating that there is no appre-
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dimerization region of protein and fluorescence studies of the mutant mini TyrRS form.

The computational models of the spatial structure of the catalytic modules were constructed on the basis of the X-ray crystallo-
graphic data of the N-terminal catalytic module of human tyrosyl-tRNA synthetase [25] and the computational model of the structure of the full-length \textit{Bos taurus} tyrosyl-tRNA synthetase [26]. The models of spatial structure of native and mutant mini \textit{Bt}TyrRS homodimers with highlighted tryptophan residues in positions 40, 87, 283 in native and in position 87 in mutant forms of the enzyme are shown in Figures 3 and 4, respectively.

Tryptophan 87 in each subunit of mini \textit{Bt}TyrRS is localized at the contact area of protein monomers of the functional mini \textit{Bt}TyrRS. The replacements of two trypto-
phan residues in mini \textit{Bt}TyrRS by alanine did not resulted in any visible changes of its 3D structure.

Similar data were obtained from the comparison of the spatial structural organization of the contact regions of protein monomers in the environment of the Trp87 residue in native and mutant forms of mini \textit{Bt}TyrRS (Fig. 5). They also did not show any obvious changes after mutagenesis in the structure of the recom-
binant enzyme.

With the help of UCSF Chimera software the environments of Trp87 residue in the sphere of radius 5 Å in both the native and mutant forms of \textit{Bt}TyrRS were visualized and analyzed (Table 1). The analysis showed that there are 11 residues in the given region around Trp87: 6 hydrophobic (Tyr79, Ala85, Leu89, Leu90, Thr121, Leu131), two negatively charged res-

![Fig. 2. Electrophoretic analysis of the expression of the mutant form of \textit{Bos taurus} tyrosyl-tRNA synthetase in \textit{E. coli} BL21 (DE3) pLysE strain and electrophoretic control of the purity of the mutant protein after chromatographic purification on Ni-NTA agarose (12 % acryl-
amide gel). M — Mixture of marker protein (“ Invitrogen “, Mark 12 Unstained Standart). 1 — Cell lysate of bacterial culture of \textit{E. coli} strain BL21 (DE3) pLysE, transformed with plasmid pET30a-39KYRSW87 before IPTG induction of mutant mini \textit{Bt}TyrRS synthesis. 2 — Cell lysate of bacterial culture of \textit{E. coli} strain BL21 (DE3) pLysE, transformed with plasmid pET30a-39KYRSW40 after IPTG induction of mutant form mini \textit{Bt}TyrRS synthesis. 3 — Mutant mini \textit{Bt}TyrRS after chromatographic purifi-
cation on Ni-NTA agarose.](image)
residues (Lys127, Arg135) and neutral Pro86. Table 1 represents the distances from the Cα atom of Trp87 to the Cα atom of the corresponding residue in the native and mutant forms of BrTyrRS. It can be seen that the Trp87 microenvironment is similar in both forms. We observed only minor changes in the solvent accessibility of Trp87.

The fluorescence spectrum of the mutant form of the catalytic modulus of Bos taurus tyrosyl-tRNA synthetase at the excitation wavelengths of 280 nm and 295 nm are shown in Fig. 6. The determined fluorescence characteristics of the mutant protein, in particular, the position of the fluorescence maximum, λm, and the half-width of the fluorescence spectra, Δλ, are 338 nm and 60 nm, respectively.

According to the three spectral classes model of tryptophan residues in protein structure, the tryptophan residue at position 87 refers to the spectral class II, which is characterized by the emission of indole fluorophore immobilized in the concavity on the surface of the protein, and does not contact with free but only
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with bound water and other polar groups in the protein structure [21].

The parameters of the fluorescence spectra of tryptophan residues depend on the polarity of their microenvironment, as well as the ability of the tryptophan residue to relax during the fluorescence lifetime. It should be kept in mind that the polarity of the tryptophan residue microenvironment is determined not only by its accessibility to the solvent molecules, but also by its own polar protein groups, which are the parts of the microenvironment [21, 22].

Previously, we have studied the intrinsic tryptophan fluorescence of native mini TyrRS and analyzed its intramolecular dynamics by fluorescence spectroscopy [27]. The analysis of the localization and microenvironments of three tryptophan residues responsible for the intrinsic fluorescence of mini TyrRS allowed us to characterize their accessibility in the structure of the enzyme dimer and the microenvironment conformational mobility. The characteristics of the tryptophan fluorescence of mutant mini TyrRS with a single tryptophan residue at position 87 indicate the immobilization of the tryptophan residue microenvironment at the dimer interface.

The mutant form of mini-TyrRS with tryptophan-87 residue, which is localized in the

| Table 1. Amino acid environment and solvent accessibility of Trp87 in native and mutant forms of mini \textit{Bt}TyrRS |
|---------------------------------------------------------------|
| **Amino acid environment of Trp87 in \textit{Bt}TyrRS (native form)** | **Distance between atoms, Å** | **Accessibility to solvent (Surface area, %)** | **Amino acid environment of Trp87 in \textit{Bt}TyrRS (mutant form)** | **Distance between atoms, Å** | **Accessibility to solvent (Surface area, %)** |
| **Tyr79A** | 4.2 | 15.6 | **Tyr79A** | 4.6 | 18.6 |
| **Ala85A** | 4.0 | 31.7 | **Ala85A** | 4.1 | 26.0 |
| **Pro86A** | 3.2 | 72.8 | **Pro86A** | 3.4 | 75.4 |
| **Glu88A** | 2.8 | 85.5 | **Glu88A** | 2.1 | 82.8 |
| **Leu89A** | 4.8 | 26.8 | **Leu89A** | 4.7 | 26.7 |
| **Leu90A** | 3.6 | 7.2 | **Leu90A** | 4.2 | 6.8 |
| **Glu91A** | 4.0 | 29.9 | **Glu91A** | 3.7 | 28.9 |
| **Thr121A** | 4.5 | 34.0 | **Thr121A** | 6.4 | 37.5 |
| **Lys127B** | 4.4 | 75.1 | **Lys127B** | 4.9 | 75.3 |
| **Leu131B** | 4.2 | 16.0 | **Leu131B** | 4.4 | 18.4 |
| **Arg135B** | 4.8 | 27.6 | **Arg135B** | 4.8 | 27.6 |

**Fig. 6.** Fluorescence spectra of the mutant form of catalytic module of \textit{Bos taurus} tyrosyl-tRNA synthetase at excitation wavelengths of 280 nm and 295 nm.
region of dimerization of the enzyme, can be effectively used to investigate the conformational changes of tyrosyl-tRNA synthetase associated with the neurodegenerative disease of Charcot-Marie-Tooth neuropathy [28, 29].

Conclusions

In this work it has been found that the replacement of Trp40 and Trp283 residues by alanine in Bos taurus mini-tyrosyl-tRNA synthetase cloned in the expression plasmid pET30a-39KYRSW87 does not affect the synthesis of the mutant form of the enzyme. The amount of soluble form of recombinant mutant mini Bt Tyrs expressed in E. coli strain BL21 (DE3) pLysE significantly increased when the temperature of incubation of bacterial culture was reduced from 37 °C to 25 °C. The yield of purified homogeneous mutant mini Bt Tyrs upon incubation of the culture E. coli strain BL21 (DE3) pLysE at 25 °C transformed with plasmid pET30a-39KYRSW87 is about 30 mg per 1L of LB culture medium. According to the fluorescence spectroscopy data and computational modeling of mini Bt Tyrs the microenvironment of Trp87 residue in mutant form is rigid at the interface of enzyme subunits.

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Виділення та характеристика мутантної форми N-кінцевого каталітичного модуля тирозил-тРНК синтетази Bos taurus з заміною Trp 40 та Trp 283 на аланин

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Мета. Отримання мутантної однотриптофанової форми N-кінцевого каталітичного модуля тирозил-тРНК синтетази Bos taurus (міні TyrRS) для моніторингу локальних конформаційних змін методом флуоресцентної спектроскопії. Методи. Біохімічна експрясія, метал-хелатуюча хроматографія, гель-електрофорез, флуоресцентна спектроскопія, комп’ютерне моделювання. Результати. Проведено заміну двох триптофанових кодонів кодонами аланина в кДНК, що кодує міні BrTyrRS. Ці мутації не впливають на синтез та розчинність міні BrTyrRS в штамі E. coli BL21 (DE3)
pLysE. Количество растворимой формы мутантной ми
BtTyrRS в цитоплазме бактериальных клеток при інку
бації бактеріальної культури при 25 °C становило
близько 47 % від загальної кількості рекомбінантного
білка. Комп’ютерне моделювання та флуоресцентное
dослідження однотриптофанової форми міні BtTyrRS
показало, що залишок Trp 87 локалізований в області
dимеризації субодиниці ферменту. Характеристики
флуоресценції триптофану мутантного міні BtTyrRS
вказують на те, що Trp 87 локалізований в іммобілізо
ваному мікросередовищі інтерфейсу димера.
Прийоми. Встановлено оптимальні умови бактері
альної експресії мутантної форми міні BtTyrRS, що
містить Trp 87 в культурі бактерій E. coli штаму BL21
(DE3) pLysE. Тrp 87-вмісна форма міні BtTyrRS може
бути використана для моніторингу локальних конфор
маційних змін в інтерфейсі димера фермента.
Ключові слова: тирозил-тРНК синтетаза, міні
TyrRS, бактеріальна експресія, флуоресцентна спектро
скопія, комп’ютерне моделювання
Выделение и характеристика мутантной
формы N-концевого каталитического модуля
tирозил-тРНК синтетазы B. taurus с заменой
Trp 40 и Trp 283 на аланин
В. Н. Засец, Д. М. Ложко, А. Ю. Цуварев,
Л. А. Коломиец, П. Е. Зуб, А. И. Корнелюк
Цель. Получение мутантной однотриптофановой
формы N-концевого каталитического модуля тиро
зил-тРНК синтетазы Bos taurus (мини TyrRS) для мо
ниторинга локальных конформационных изменений
методом флуоресцентной спектроскопии. Методы.
Бактериальная экспрессия, металл-хелатирующая
хроматография, гель-электрофорез, флуоресцентная
спектроскопия, компьютерное моделирование.
Результаты. Проведена замена двух триптофановых
кодонов кодонами аланина в kДНК, кодирующей мини
BtTyrRS. Эти мутации не влияют на синтез и раство
римость мини BtTyrRS в штамме E. coli BL21 (DE3)
pLysE. Количество растворимой формы мутантной
мини BtTyrRS в цитоплазме бактериальных клеток при
инкубации бактериальной культуры при 25 °C соста
вило около 47 % от общего количества рекомбинант
ного белка. Компьютерное моделирование и флуорес
ценное исследование однотриптофановой формы
мини BtTyrRS показало, что остаток Trp 87 локалізо
ван в области димеризации субъединицы фермента.
Характеристики флуоресценции триптофана мутант
ной мини BtTyrRS указывают на то, что Trp 87 лока
лизован в иммобилизованной микросреде интерфейса
dимера. Выводы. Установлены оптимальные условия
бактериальной экспрессии мутантной формы мини
BtTyrRS, содержащий Trp 87, в культуре бактерий
E. coli штамма BL21 (DE3) pLysE. Trp 87-содержащая
форма мини BtTyrRS может быть использована для
мониторинга локальных конформационных изменений
в интерфейсе димера фермента.
Ключевые слова: тирозил-тРНК синтетазы,
мини TyrRS, бактериальная экспрессия, флуоресцентная
спектроскопия, компьютерное моделирование

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