INTRODUCTION

tRNA-guanine transglycosylase (TGT) catalyzes the replacement of guanine with queuine in human tRNA and participating in the translation mechanism, is involved in the development of cancer. However, information on the small-molecule inhibitors that can suppress activity of this enzyme is very limited. Molecular dynamics simulations were used to determine the amino acid residues that provide efficient binding of inhibitors in the active site of tRNA-guanine transglycosylase. It was demonstrated using 7-methylguanine molecule as a probe that the ability of the inhibitor to adopt a charged state in the environment of hydrogen bond acceptors Asp105 and Asp159 plays a key role in complex formation. Formation of the hydrogen bonds and hydrophobic contacts with Gln202, G1y229, Phe109, and Met259 residues are also important. It has been predicted that introduction of the substrates would have a different effect on the ability to inhibit tRNA-guanine transglycosylase, as well as the DNA repair protein poly(ADP-ribose) polymerase 1, which can contribute to the development of more efficient and selective compounds.

**Abbreviations**: 7-MG, 7-methylguanine; MD, molecular dynamics; PARP-1, poly(ADP-ribose) polymerase 1; TGT, tRNA-guanine transglycosylase.

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tumors in the xenograft model. It should also be mentioned that enhanced expression of TGT was found in lung cancer patients, and the expression level correlated with survival [12, 13]. Involvement of the human TGT in pathological processes indicates the need for the development and testing of selective inhibitors of this protein.

Inhibition of human TGT by low-molecular-weight compounds has been poorly investigated. The main source of information about potential inhibitors is publication devoted to testing structural analogues of the substrate against TGT from rabbit reticulocytes [14]. 7-Methylguanine (7-MG, natural nitrogenous base) was found to be one of the most effective inhibitors, competitive with respect to queuine (Fig. 1, \(K_i = 1 \mu\text{M}\)) [14, 15]. Some other studies have shown that 7-MG is also able to suppress the activity of bacterial TGT [16, 17]. Alignment of TGT sequences from different organisms, revealing conserved active site residues, is shown in Online Resource 1, Fig. S1. Interestingly, 7-MG suppresses activity of the human poly(ADP-ribose) polymerase 1 (PARP-1, \(K_i = 60 \mu\text{M}\)) [18-21] which plays an important role in pathologies of cardiovascular, nervous, and respiratory systems, as well as in cancer diseases [22-25]. A more selective effect on the target may be required for clinical use of TGT inhibitors, and therefore, structural optimization of 7-MG and its analogues is of great interest.

In the present work, interaction of the human TGT with the 7-MG inhibitor was modeled, and recommendations were given for further development of more effective and selective inhibitors.

**MATERIALS AND METHODS**

For molecular mechanical description of 7-MG in a cationic state, parameters of the Amber-compatible field ff99 [26] were used with exception of partial charges. Assignment of partial atomic charges was carried out by the RESP method (see in Online Resource 1, Fig. S2) [27, 28]. For this purpose, molecular electrostatic potential was preliminarily calculated at the HF/6-31G* level of theory using GAMESS [29]. Parameters of the uncharged 7-MG molecule were obtained in a similar way in our previous work [18]. Amino acid residues of the protein were described by the ff14SB force field with improved torsion parameters [30].
Molecular model of the human TGT was constructed based on the crystal structure of an enzyme–substrate complex containing a queuine molecule (PDB ID 6h45, chain A) [4]. Coordinates of the Arg68-His93 and Asn101-Asp105 regions were transferred from the 7nq4 structure [31]. Coordinates of the 7-MG inhibitor in the active site were obtained by superimposing the fused rings of 7-MG and queuine. Next, the structure of the enzyme-inhibitor complex was optimized and studied by molecular dynamics (MD) method using AmberTools 20 and Amber 20 [32-34]. Hydrogen atoms were added taking into account ionization properties of the amino acid residues; when protonating the histidine side chains, their molecular environment was considered. The structure was placed in a truncated octahedral TIP3P water box (minimum distance from the protein to the box edge was 12 Å); Cl− ions were added to neutralize the total charge.

At the first stage of energy minimization of the obtained system (2500 steepest descent steps + 2500 conjugate gradient steps), the protein and 7-MG coordinates were kept fixed by positional restraints of 2 kcal/(mol Å²) on heavy atoms. The second stage of minimization (5000 steepest descent steps + 5000 conjugate gradient steps) was performed without any restraints. The system was then heated up from 0 to 300 K using positional restraints of 1 kcal/(mol Å²) on the protein and 7-MG atoms (250 ps, constant volume) and equilibrated at 300 K (25 ns, constant pressure). Acquisition of the equilibrium conformation of the complex was confirmed by analyzing the root-mean-square deviation of the protein backbone atoms from the initial position (mobile loops and terminal residues of the protein were not taken into account). A 100 ns trajectory of the equilibrium MD simulation was then calculated and analyzed. More detailed simulation protocols are given in Online Resource 2, Table S1. The calculations were carried out on the supercomputer “Lomonosov-2” using graphics processing units (Nvidia Tesla K40M).

Docking of the 7-MG molecule into the TGT active site was carried out using the Lead Finder 1708 program in the “extra precision” mode [35, 36]; an energy grid map covered the binding site of the queuine molecule. The structures were visualized using the VMD 1.9 program [37].

RESULTS AND DISCUSSION

The human TGT model was constructed on the basis of the 6h45 crystal structure that describes interaction of the QTRT1 catalytic subunit with the queuine substrate. In general, this complex with a resolution of 2.4 Å was appropriate for preparing the starting model, however, it was decided to transfer some parts of the protein chain from the 7nq4 structure obtained by the same authors. The first region, Arg68-His93, is less ordered in the 6h45 structure compared to other TGT structures, which may be related to crystallization conditions. Through a network of hydrogen bonds, the Arg68-His93 residues affect conformation of another region, Asn101-Asp105, which was also transferred from the 7nq4 structure.

In modeling of the 7-MG binding to the TGT active site ionization properties of this inhibitor were taken into account. Previously, interaction of the structurally similar compounds with bacterial TGT was studied using isothermal calorimetry, site-directed mutagenesis, and X-ray crystallography. It was suggested that the presence of two negatively charged aspartate residues, Asp105 and Asp159 (human TGT numbering), promotes transition of the inhibitors to cationic form upon binding in the active site [38, 39]. The corresponding transition of 7-MG (protonation on the N3 atom) is shown in Fig. 2. The positively charged inhibitor is able to interact more efficiently with Asp105 and Asp159, so it was decided to model 7-MG in both cationic form and neutral form (as a control).

7-MG is a competitive inhibitor of TGT and a structural analogue of the queuine substrate, so it is expected to bind in a similar manner. The starting position of 7-MG in the active site of the TGT model corresponded to coordinates of the 7-deazaguanine fragment of queuine. The enzyme–inhibitor complex was then equilibrated and subjected to a 100 ns MD simulation to generate a trajectory, the analysis of which made it possible to characterize intermolecular interactions (table, Fig. 3). First
of all, formation of the hydrogen bonds of 7-MG with Asp159 and Asp105 should be mentioned, since interaction of the delocalized positive charge of the inhibitor with the negatively charged carboxyl groups could make a significant contribution to the binding energy. In that case one of the bonds with Asp105 is mediated by the protonated nitrogen atom N3. Additional hydrogen bonds are formed with the Gln202 and Gly229 residues. Purine rings of 7-MG form \(\pi\)-stacking with Phe109, as well as hydrophobic contact with the side chain of Met259. The methyl group of 7-MG is in contact with the benzene ring of Phe109.

Modeling of the TGT complex formation with the neutral form of 7-MG showed a significant decrease in the stability of interactions with the residues of the active site compared to its cationic form. Hydrogen bonds with the Asp159 and Gly229 residues were episodic, which affected the average distance between atoms, and bonds with Asp105 and Gln202 were actually lost (see Online Resource 2, Table S2). The side chain of Asp105 was found to be turned away from the inhibitor molecule (Online Resource 1, Fig. S3 and S4), which is apparently due to the absence of a proton on the N3 atom. It is noteworthy that the related compound 3-deazaguanine, in which the N3 atom is replaced by carbon (Fig. 1), has an inhibitory activity two orders of magnitude lower compared to 7-MG [14]. Thus, we can conclude that the effective binding of 7-MG molecule in the active site of TGT is possible due to its transition to a cationic form.

Since 7-MG also inhibits the DNA repair protein PARP-1, which is not similar in sequence to TGT, it is necessary to reveal structural factors that determine the possibility of its binding to two different targets, as well as possible ways of inhibitor modifications for a more selective action. Previously, it was experimentally shown that the important factor for effective binding of purine derivatives and analogues with TGT and PARP-1 is the presence of amino group at the position 2 and oxo group at the position 6 [14, 18]. In the case of TGT, the amino group forms the above-mentioned hydrogen bonds with
Asp105 and Asp159 (Fig. 3), and in the case of PARP-1, it interacts with the key residue Gly863 (Fig. 4). The oxo group forms hydrogen bonds with the Gln202 and Gly229 residues of TGT and with the Gly863 and Ser904 residues of PARP-1. Special attention should be paid to high similarity of the \( \pi \)-stacking interaction: in the case of TGT, it is mediated by the Phe109 side chain, and in the case of PARP-1, by the Tyr907 side chain. Taken together, these interactions facilitate binding of the inhibitors to both TGT and PARP-1.

A possible way to optimize the structure of 7-MG and its analogues is to modify amino group at the position 2 or methyl group at the position 7. The first option is suitable for the PARP-1 inhibitors, since the introduced substituent is oriented towards the additional binding site of the NAD\(^+\) adenine group (Fig. 4a). In the case of TGT, the amino group forms hydrogen bonds with Asp105 and Asp159, therefore, any modification of this group will disrupt these interactions [40]. The second option (modification of the methyl group), on the contrary, is not suitable for PARP-1, since the corresponding binding pocket formed with participation of the Ala898 residue (Fig. 4b) is very limited in size. In the case of TGT, the 7-MG methyl group is oriented towards the binding site of cyclopentene moiety of the queuine substrate formed by the residues Val161-Ser164 and Gly229-Gly232. This site is able to interact with various groups of atoms, which is confirmed by the broad substrate specificity of human TGT, demonstrated using synthetic derivatives of 7-deazaguanine [41].

The molecular model of human TGT enables searching for inhibitors by docking and virtual screening approaches. The most promising compounds can be selected from a large library by evaluating binding energies and analyzing interactions with the crucial amino acid residues [42-44]. As an illustrative example, we performed docking of the 7-MG molecule against the ensemble of TGT structures obtained by MD simulation. For this purpose, structures were extracted at 10 ns intervals over the 100 ns trajectory of the equilibrium TGT simulation, and water and inhibitor molecules were removed from them. Next, 7-MG was docked into the region covering the binding site of the queuine substrate, and thus the ensemble of enzyme–inhibitor complexes was obtained (Online Resource 1, Fig. S5). Analysis of the obtained models showed that the docking well reproduced interactions of 7-MG with the amino acid residues of TGT described above. The ensemble average value of the binding free energy \( \Delta G^\text{calc} \) was \(-7.4\) kcal/mol. According to the formula \( \Delta G = RT \ln(K_i) \), this energy value corresponds to the inhibition constant of 4 \( \mu \)M, which is in good agreement with the experimental data (\( K_i = 1 \mu \)M) [14].

The conducted study made it possible to draw some conclusions that are important for designing inhibitors of human TGT, a promising target for the treatment of oncological diseases. (i) The inhibitor may adopt a charged state upon binding in the active site in the environment of Asp105 and Asp159, which leads to the formation of more stable hydrogen bonds with these residues. This feature is important for membrane trans-
port, as inhibitor molecule becomes positively charged only upon binding to the target protein inside the cell. (ii) The possibility of hydrogen bonding and hydrophobic contacts with Gln202, Gly229, Phe109, and Met259 residues should also be taken into account in the inhibitor design. (iii) A more selective interaction of the inhibitor with TGT could be achieved by elongation of the structure towards the binding site of the substrate’s cyclopentenone group formed by the Val161-Ser164 and Gly229-Gly232 residues.

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