ALANINE SCANNING OF DINITROANILINE/PHOSPHOROTHIOAMIDATE SITE OF α-TUBULIN IN PLASMODIUM SPECIES DISTRIBUTED IN INDIA

Aim. Identification of amino acid residues participating in specific binding of dinitroaniline and phosphorothioamidate compounds with α-tubulin in Plasmodium falciparum. Methods. Protein structure modelling, protein structure optimization using molecular dynamics method, ligand-protein docking, alanine scanning mutagenesis. Results. Molecular docking of canonical compounds and alanine scanning mutagenesis, indicate two key (Arg2, Val250) and one minor (Glu3) residues involved in binding of both - dinitroaniline and phosphorothioamidate compounds. At the same time, it was revealed two minor residues (Asp251, Glu254) interacting only with some members of dinitroaniline grope. Conclusions. It was identified amino acid residues predetermining existence of joint site and similar interaction of α-tubulin with dinitroaniline and phosphorothioamidate compounds in P. falciparum.

Keywords: malaria, Plasmodium, α-tubulin, molecular interaction, dinitroanilines compounds, phosphorothioamidate compounds, alanine scanning mutagenesis.

Human malaria is a complex disease caused by numerous Plasmodium species, which threatens the half of Earth population. The control of malaria infection is hampered by many factors, including emerging of drug resistance. It is a fact that many of existing malaria therapeutics are increasingly ineffective and it is an urgent need in development of principally new therapeutic strategies and agents [1]. It should be noted that malaria is an important component of morbidity and mortality in the Republic of India. The National Vector Borne Disease Control Program of India reported ~1.6 million cases and ~1100 malaria deaths in 2009 [2].

Malaria in India is known to be caused at least by four Plasmodium species: P. falciparum, P. vivax, P. ovale and P. malariae [3]. Two of them, P. falciparum and P. vivax, are dominative in this area. P. malariae has been reported in the eastern India state of Orissa [4], while P. ovale appears to be extremely rare if not absent [2]. Among the aforementioned species P. falciparum is most severe strain of the malaria species correlated with almost every malarial death (CDC - https://www.cdc.gov/malaria/about/disease.html). P. falciparum is strongly associated with severe disease syndrome known as cerebral malaria, which is associated with high mortality [2]. At the same time, it is known about the existence of different ecotypes and lines in both P. falciparum and its mosquito vector [3].

A number of drugs have been developed to treat malaria. However, with emergence of resistance, many of the previously effective substances have lost their relevance (quinine, chloroquine, amodiaquin, pyrimethamine, etc.). Due to point mutations, Plasmodium increased resistance against complex drugs of antifolate type, of which sulfadoxine and pyrimethamine were the most commonly used [5].

During past decade years there has been a new growth of interest in tubulin as an important target for compounds with antiprotozoan activity. Several classes of microtubule (MT) inhibitors have demonstrated potent activity against malarial parasites in in vivo: vinblastine [6 – 8], dolastatin 10 [9], auristatins [10] and taxoids [11, 12]. Most of these agents have been demonstrated to disrupt or stabilize normal microtubular structures. Unfortunately, most all these compounds show toxicity to mammalian cells [10, 13] due to the interspecies conservation of tubulin [14]. P. falciparum and human α-tubulins share ~83% and β-tubulins ~87% of identity. However it was found that human antimalarial drugs (e.g. sulfadiazine, sulfadoxine, pyrimethamine, cycloguanyl) were lethal for the model plant Arabidopsis thaliana at similar
concentrations to commercial herbicides: glufosinate and glyphosate [15]. Although MT inhibitors from anticancer programs have proved useful in probing MT function in parasites, such non-selective agents have no prospects as antimalarial drugs [13, 14].

Previous studies have shown that dinitroaniline and phosphorothioamidates compounds, which are active against plant microtubules, are also active against P. falciparum, and may act as antimalarial drugs [6, 15]. It indicates α-tubulin as extremely promising molecular target in the case of parasitic diseases such as leishmaniasis and trypanosomiasis [16, 17]. In plants, phosphorothioamidates demonstrate effects, similar to dinitroanilines and bind to the same binding site. Phosphorothioamidates have similar effects on plants as the dinitroanilines and bind to the same molecular site. As phosphorothioamidates have a more than 100-fold higher solubility in aqueous solutions than dinitroanilines, these compounds are more promising candidates for modification than dinitroanilines, where biological studies indicated that maintaining high sufficient drug concentrations is difficult [18]. Therefore we consider dinitroaniline and phosphorothioamidate compounds as the most priority group for the search of new antimalarial agents.

The purpose of current study was application of alanine scanning mutagenesis for identification of amino acids playing key role in binding of dinitroaniline and phosphorothioamidates compounds with Plasmodium α-tubulin.

Materials and methods

Structural model of α-tubulin molecule from Plasmodium falciparum (TBA_PLAFK, UniProtKB: P14642) was built using protein structure homology-modelling server Swiss-Model [19]. The template modelling was based on template RCSB Protein Data Bank (www.rcsb.org) [19] structures: 5UBQ (A) - α-tubulin from cilia of Tetrahymena thermophila (Cryo-EM structure) [21] and 2.5 E X-RAY structure 5KX5 (A) of α-tubulin from Ovis aries [22].

For protein structure geometry optimization we used minimization of potential energy based on steepest descent algorithm with a maximum number of steps = 1000 and a gradient = 0.1 of charmm27 force field. All visualizations and analysis of PDB-structures and constructed model of Plasmodium α-tubulin were performed using PyMOL v.1.5.0.5 software (www.pymol.org).

Alanine scanning mutagenesis was performed using VMD/NAMD software, enhanced with AlaScan plugin (Version 1.0) [23]. As the ligands we used 3D-models of 18 dinitroaniline and 4 phosphorothioamidates compounds.

Dinitroaniline compounds: Sulfamidas16 (PubChem CID: 11282001), Sulfamidas21 (PubChem CID: 11235040), Sulfamidas20 (PubChem CID: 11199668), Sulfamidas23 (PubChem CID: 11177910), Sulfamidas33 (PubChem CID: 11177051), Sulfamidas24 (PubChem CID: 11165550), Sulfamidas25 (PubChem CID: 11155360), Benzenesulfonamide (PubChem CID: 10428592), CHEMBL80689 (PubChem CID: 10250523), CHEMBL78502 (PubChem CID: 10193905), Ethalfluralin (PubChem CID: 41381), Isopropaline (PubChem CID: 36606), Fluchloralin (PubChem CID: 36392), Prodimine (PubChem CID: 34469), Dinitramide (PubChem CID: 34468), Profuralin (PubChem CID: 33500), Diporalin (PubChem CID: 15966), Benfluralin (PubChem CID: 2319); and

Phosphorothioamide compounds: CHEMBL1835180 (PubChem CID: 56669570), CHEMBL1835273 (PubChem CID: 56659918), CHEMBL1835163 (PubChem CID: 14179764), Amiprophos (PubChem CID: 36612).

Results and discussion

Since our main task is the search for new effective inhibitors of Plasmodium α-tubulin, it was necessary to identify key amino acids of joint dinitroaniline/phosphorothioamide site that playing a key role in the ligand binding. One of the most common methods for such selection of amino acids important for protein-protein or protein-ligand interaction is the method of alanine scanning mutagenesis [24].

Alanine (Ala) scanning is a widely used mutagenesis approach in which residues in a target protein are systematically substituted for alanine at selected positions by site-directed mutagenesis (in silico as well as in genetical experiment). Substitution with alanine residues eliminates side-chain interactions without altering main-chain conformation or introducing steric or electrostatic effects, so is often the preferred choice for testing the contribution of specific side-chains while preserving native protein structure. In most cases, replacement of the native amino acid(s) with an alanine residue(s) does not change overall conformation of polypeptide chain, such as in the opposite cases of glycine or proline substitutions. Also, such replacement
with alanine never is accompanied by electrostatic or steric effects. It is also known that alanine is very common in both internal and external regions of protein globules. Thus, such virtual “mutations”, i.e. scanning with neutral alanine, make it possible to identify key amino acids important for enzyme active center, protein activity, or participating in protein-protein and protein-ligand interactions. Thus, the alanine scanning allows us to investigate the structural and functional aspects of protein-ligand and protein-protein interactions [25, 26].

In current study, we used the full-atom model of α-tubulin from P. falciparum (TBA_PLAFK, UniProtKB: P14642), and the library of canonical dinitroaniline/phosphorothioamidate ligands (22 compounds: trifluralin-, orizalin- and amiprophosmethyl-like compounds).

Initial ligand structures were obtained from the PubChem Compound database through searching of substances similar to known tubulin inhibitors of dinitroanilines/phosphorothioamidate group. Similar structures were selected based on Tanimoto Threshold (TT) in 90-95% (in the case of trifluralin- and orizalin-like compounds).

Table. Summary results of alanine scanning effect on the binding of dinitroaniline and phosphorothioamide to the surface of α-tubulin from Plasmodium falciparum

| Grop of compounds | CID      | ARG2 | GLU3 | GLN133 | ARG243 | VAL250 | ASP251 | GLU254# |
|------------------|----------|------|------|--------|--------|--------|--------|---------|
| Amiprophosmethyl-like | 5669570  | $$$  |      |        |        |        |        |         |
|                   | 56659918 | $$$  |      |        |        |        |        |         |
|                   | 14179764 | **** | *    |        |        |        |        |         |
|                   | 36612    | *****| *    | ****   |        |        |        |         |
| Orizalin-like     | 11282001 | *    | *    | ****   |        |        |        |         |
|                   | 11235040 | *    | *    | ****   |        |        |        |         |
|                   | 11199668 | **   | **   | ***    | *      |        |        |         |
|                   | 11177910 | *****| **   | ****   |        |        |        |         |
|                   | 11177051 | **   | **   | ****   |        |        |        |         |
|                   | 11165550 | **** | *    | ****   |        |        |        |         |
|                   | 11155360 | *    | *    | ****   |        |        |        |         |
|                   | 10428592 | *    | *    | ****   |        |        |        |         |
|                   | 10250523 | **** | *    | ****   |        |        |        |         |
|                   | 10193905 | **** | *    |        |        |        |        |         |
| Trifluralin-like  | 41381    | *    | *    | ****   |        |        |        |         |
|                   | 36606    | ***  | *    | ****   |        |        |        |         |
|                   | 36392    | *    | *    | ****   |        |        |        |         |
|                   | 34469    | **   | *    | ****   |        |        |        |         |
|                   | 34468    | *    | *    | ****   |        |        |        |         |
|                   | 33500    | **   | *    | ****   |        |        |        |         |
|                   | 15966    | **   | *    | ****   |        |        |        |         |
|                   | 2319     | **   | *    | ****   |        |        |        |         |

Notes: importance for binding by growth: $$>****>***>**>*>****>*****.
Conclusions
These studies identify amino acid residues and interactions, predetermining existence of joint site and similar interaction of α-tubulin with dinitroaniline and phosphorothioamide compounds in *Plasmodium falciparum*. Alanine scanning mutagenesis indicate two key (Arg2, Val250) and one minor (Glu3) residues involved in binding of both - dinitroaniline and phosphorothioamide compounds. At the same time, it was revealed two minor residues (Asp251, Glu254) interacting only with some members of dinitroaniline group. Despite existence of the general mechanism of dinitroaniline and phosphorothioamide binding, alternative interactions within the previously defined site are enough realistic. We assume that these differences can contribute total binding energy and predetermine variations in binding of studied compounds in the site. Our data indicate that in the case of dinitroaniline compounds such differences may be stronger than in the case of phosphorothioamide.

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