INTRODUCTION

Malaria caused by the parasite *Plasmodium falciparum* is one of the major devastating diseases worldwide, with half a billion cases and up to 0.5-1 million deaths, mostly among young children, yearly. According to the World Health Organization, most deaths result from severe malaria (SM) complications, the major forms of which include cerebral malaria (CM), placental malaria (PM), respiratory distress, and severe anemia [1].

Currently, there is a noticeable decline in the efficacy of commonly used anti-malarial drugs. It should be noted that although still efficient anti-malarial drugs rapidly kill the parasite, significant mortality (10-15%) ensues in SM, particularly within 24 h of hospital admission. This happens most likely due to the cytoadherence of infected erythrocytes (IEs) that continues long after the parasites inside the red blood cells have been killed [2]. Unfortunately, novel parasite-killing drugs will have the same problem if they do not affect the cytoadhesion of IEs. This strongly justifies the need for the development of adjunctive therapies able to prevent and/or reverse the pathophysiological consequences of cytoadherence in SM. Our main objective was to discover, characterize, and optimize novel inhibitors of cytoadhesion of *P. falciparum*-infected erythrocytes to the host vasculature for the treatment of severe forms of malaria (including CM and PM). Below, we review our works in this area.

ADVANTAGES OF ANTI-ADHESION DRUGS IN SEVERE MALARIA

Most anti-malarial drugs target intracellular processes required for the parasite survival and propagation. Such drugs must penetrate into the parasite by crossing several membranes, including erythrocyte plasma mem-

**Abbreviations**: CM, cerebral malaria; DMSO, dimethyl sulfoxide; HT, high-throughput; IEs, erythrocytes infected with *Plasmodium falciparum*; PM, placental malaria; SM, severe malaria.

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**Malaria Parasite *Plasmodium falciparum* Proteins on the Surface of Infected Erythrocytes as Targets for Novel Drug Discovery**

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**Abstract**—Specific adhesion (sequestration) of *Plasmodium falciparum* parasite-infected erythrocytes (IEs) in deep vascular beds can cause severe complications resulting in death. This review describes our work on the discovery, characterization, and optimization of novel inhibitors that specifically prevent adhesion of IEs to the host vasculature during severe malaria, especially its placental and cerebral forms. The main idea of using anti-adhesion drugs in severe malaria is to release sequestered parasites (or prevent additional sequestration) as quickly as possible. This may significantly improve the outcomes for patients with severe malaria by decreasing local and systemic inflammation associated with the disease and reestablishing the microvascular blood flow. To identify anti-malarial adhesion-inhibiting molecules, we have developed a high-throughput (HT) screening approach and found a number of promising leads that can be further developed into anti-adhesion drugs providing an efficient adjunct therapy against severe forms of malaria.

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**In memory of Lev Pavlovich Ovchinnikov, a good friend and a great scientist**
brane, parasitophorous vacuolar membrane, and parasite plasma membrane. These drugs also have potential side effects, since they may similarly penetrate into the host cells and interfere with the host cellular functions.

Cytoadherence plays an important role in the lifecycle and virulence of *P. falciparum* (reviewed in [3]), the deadliest of human malaria parasites, and is a cause of PM [4] and CM [5-7] (figure). It is reasonable to suggest that reducing cytoadherence of IEs to the host vasculature will have a strong beneficial effect on the host defense against the infection, a hypothesis that can only be tested experimentally. A simple fact that the appearance of anti-adhesion antibodies preventing IE cytoadhesion to the placental syncytiotrophoblast in multigravid women is associated with protection against PM pathology [8, 9] indicates that the anti-adhesion therapy might be effective against at least this severe disease (figure). The anti-adhesion activity is considered as an important factor in the development of successful vaccine against PM [10]. Furthermore, we found that >75% immune adults have high levels of antibodies inhibiting adhesion to the ICAM-1 receptor [11] involved in SM and CM (see below). This anti-adhesion activity may contribute (along with the ability to opsonize IEs) to the fast recovery of severely sick children injected with IgG from immune adults [12]. The time window for the pharmacological effect of drugs is extremely critical for the survival in SM. The anti-adhesion agents working from outside of IEs, will likely have a higher speed of action. In this respect, molecules with a limited ability to penetrate through the plasma membranes and, therefore, to interfere with the host functions, might be especially safe, although they may need to be delivered intravenously, which is well justified in the case of life-threatening SM episodes. This is extremely important in the treatment of malaria during pregnancy, as the safety of both mother and fetus is the main issue with currently used drugs [13]. Moreover, as IEs bind to functionally relevant parts of host receptors and may inhibit their normal physiological functions [14], anti-adhesion drugs would have an additional positive effect on the host. In our opinion, as well as in the opinion of other researchers, anti-adhesion drugs may be very efficient and safe for the anti-malarial therapy [15, 16] when used as an adjunctive therapy together with parasiticidal drugs (figure). If, by chance, an anti-adhesion drug also has a parasiticidal effect [16], it would be even more valuable.

**CYTOADHESION OF IEs IS MEDIATED BY A FAMILY OF PfEMP1 PROTEINS**

*P. falciparum* is the only malaria parasite species that was clearly shown to efficiently sequester IEs in various organs using host cell surface receptors for adhesion to endothelial cells, uninfected red blood cells, and cells of immune system [17-21]. Cytoadhesion allows IEs to avoid clearance in the spleen (main innate anti-malarial mechanism). However, cytoadhesion also leads to the vascular occlusion that strongly correlates with disease severity [22] and inflammation of endothelial cells and various organs [23-28]. The main parasite ligand for the IE adhesion is a family (~60 members) of PfEMP1 proteins [29-31] that are expressed on the surface of IEs in specialized structures called knobs [32]. No parasite coreceptor molecules required for the IE adhesion have
been discovered so far. PfEMP1 proteins are coded by var genes. Mature parasite expresses on the surface the product of only one of the var genes [33], but can switch to another variant, thus changing its antigenic and adhesion phenotype [30, 34] (figure). Each PfEMP1 protein consists of one intracellular, one trans-membrane, and several extracellular domains (∼30-45 kDa, 2 to 7 domains per protein) heavily cross-linked by disulfide bonds [31, 35]. These domains have been classified using Greek letters for classes and numbers for sub-classes into Duffy binding-like (DBL) domains (α, β, γ, δ, ε, ζ, Χ) and cysteine-rich interdomain regions (CIDRs) (α, β, and γ) [31, 35, 36]. All DBL domains have a similar scaffold [37-40], and the C-terminal part of the CIDR scaffold is similar to the C-terminal part of the DBL domain [41]. These domains are responsible for the adhesion to various host receptors (reviewed in [42]), each domain representing a functional binding (with higher or lesser efficiency) unit. The specificity and combination of these domains in the expressed PfEMP1 may determine the final IE adhesion specificity, strength of binding, and, possibly, disease pathology [43]. We have cloned and expressed a library of PfEMP1 constructs (n > 180) that contained all domains (n = 272) or domain tandems (for example, many DBL domains are followed by the CIDR domains) from all PfEMP1 proteins (n = 59) from the NF54 parasite line, in their functional (receptor-binding) form, as has been demonstrated in a number of our publications [11, 44-49]. This proteome-wide collection of PfEMP1 constructs is an important tool for identification of various novel receptor-domain pairs and screening for anti-adhesion compounds against different binding IE specificities. Although the members of the PfEMP1 family belong to the variable class proteins (the sequences vary from parasite to parasite in the field), such variability may be not an insurmountable problem, as we discuss below.

SEVERE MALARIA SYNDROMES AND CYTOADHESION

As we mentioned above, cytoadhesion of IEs is either a cause or a prerequisite for SM, including CM [5-7, 27] and PM [4, 50]. The most studied syndrome is PM. Only one member of the PfEMP1 family, VAR2CSA, is involved in PM [51-53]. It has a characteristic sequence of three DBL-X domains followed by three DBLe domains. This architecture of VAR2CSA is conserved in all studied P. falciparum strains. Anti-adhesion antibodies, that appear after one or two successive pregnancies in malaria endemic regions, alleviate PM by inhibiting IE binding to the CSA receptor in the placenta [54]. Common drugs are not particularly effective in the prevention of PM and might be even harmful [13]. In light of this, a drug that does not penetrate cell membranes and, consequently, mother and fetus tissues, but interferes with the parasite adhesion (the key event in PM) would be particularly safe for the use in pregnant women.

As has been shown by us and other researchers, women that acquire resistance to pregnancy malaria, develop antibodies to VAR2CSA over successive pregnancies [44, 50]. Identification of a monoclonal antibody inhibiting the binding of several strains of PM parasites to the host ligand CSA indicates that the conserved features of these variable proteins are sufficient to provide a wide cross-strain effectiveness [55]. Such conserved structures of the VAR2CSA domains could be the targets for anti-adhesion compounds through the competitive or allosteric mode of action.

The main candidate for the host adhesion receptor in CM is ICAM-1 [6, 7, 27, 56]. However, it was suggested recently that PfEMP1 proteins containing a tandem of domains binding to ICAM-1 and endothelial protein C receptor (EPCR) are involved predominantly in CM pathogenesis [57, 58], potentially binding to one or both receptors [59, 60]. Only six DBLβ domains (out of more than 250 domains from ∼60 PfEMP1 proteins) in P. falciparum IT4 strain have been identified as capable of binding to ICAM-1 [61], as well as only two DBLβ domain in the 3D7 strain [11, 62]. Similarly, only a limited set of domains in each genome variant bound EPCR [63, 64]. The fact that only a small number of PfEMP1 domains strongly bind to either of these receptors fits well with the epidemiology of CM, which is a rare but extremely deadly SM syndrome. Extensive epidemiological data [65, 66] support the hypothesis that only a limited number of parasite lines, possibly expressing particular variants of PfEMP1 (and containing particular domains), may be the source of severe disease, as it happens in PM.

VASCULAR INFLAMMATION IS A COMMON RESPONSE TO IE CYTOADHESION AND A KEY EVENT IN SEVERE MALARIA

Several studies have clearly demonstrated that IE cytoadhesion directly affects vascular inflammation and endothelial dysfunction [67, 68]. Vascular inflammation contributes to the disease severity [69], plays a key role in CM and PM, and is characterized by induction of adhesion molecules (e.g., ICAM-1) and pro-inflammatory cytokines, as well as by vascular permeability caused by the disruption of the endothelial barrier integrity [70-72]. Some of these inflammatory processes are initiated and stimulated by direct interactions between PfEMP1 domains and host receptors through a complex intracellular signaling networks [25, 26]. Consequently, anti-adhesion drugs that prevent or reverse these interactions may reduce inflammation, thus contributing to a better outcome of SM syndromes. Our recent preliminary results demonstrate that the anti-adhesion treatment may reduce vascular inflammation, although further studies on this topic are required to support this observation.
VARIABILITY OF PfEMP1 FAMILY MEMBERS IN THE FIELD AS A CHALLENGING FACTOR

The sequences of PfEMP1 family members significantly vary in the field. Nevertheless, this variability does not preclude specific groups of domains from binding to the corresponding conserved receptors. This was corroborated by the recent structural study that revealed conserved binding pockets in variant PfEMP1 domains that bind to the conserved receptors [14]. In this regard, all DBL and CIDR domains have a similar scaffold [37-41]. Theoretically, molecules that mimic the binding motifs of host receptors might be effective against multiple variants of relevant PfEMP1 proteins, as demonstrated for the small molecule that mimics the ICAM-1 loop and inhibits the binding of two parasite variants to the ICAM-1 receptor [73]. However, these types of anti-adhesion molecules might be problematic, since they may compete with the natural ligands of these receptors and produce harmful side effects. Therefore, molecules that bind directly to variable PfEMP1 domains and inhibit their binding to the specific receptors would be preferable. The above-mentioned structural study [14] strongly supports the idea of inhibiting the binding of variant domains to the corresponding receptor via interaction between a single molecule and the receptor-binding conserved pocket in the protein.

Importantly, the available data support the idea that the diversity of variants involved in SM cases is also significantly restricted. In malaria endemic areas, SM accounts for <2% of total malaria cases in young children and predominantly occurs only once or twice in a lifetime [65]. This indicates that the broad immunity against SM develops quickly in early childhood [66] despite variations in the IE surface proteins. The existence of a monoclonal antibody that inhibits the binding of several pregnancy malaria parasite strains to the host ligand chondroitin sulfate A (CSA) [55] further indicates that the conserved features of PfEMP1 proteins might be sufficient to provide the cross-strain effectiveness of anti-adhesion molecules. Hence, a single adhesion-inhibiting molecule might exhibit cross-strain/PfEMP1 activity for a particular host receptor.

STRATEGY FOR IDENTIFICATION OF ANTI-ADHESION COMPOUNDS

Direct screening of live laboratory or field isolates for identification of agents that would inhibit parasite adhesion to different host receptors is an extremely slow and tedious process not currently amenable to the use of high throughput (HT) protocols due to technical and biological limitations. Our two-step approach to identification of anti-adhesion molecules overcomes these limitations [47]. Moreover, it was designed to screen mixture-based libraries to allow fast deconvolution to individual active compounds in a few steps, because the first HT step is performed in vitro. Testing the sets of such compounds for the inhibition of live parasite binding is not feasible due to the toxic effects of these mixtures or of the solvent dimethyl sulfoxide (DMSO, tolerated up to 20% in our in vitro system), as well as due to the low throughput of live parasite assays.

First, mixture-based libraries of small molecules are screened and deconvoluted to single compounds in a HT testing for the inhibition of interactions between known host receptors involved in IE adhesion and various relevant PfEMP1 domains immobilized on BioPlex beads (Bio-Rad, USA). Second, molecules identified in the first step are validated for the adhesion-inhibiting activity using live heterologous IEs. The use of heterologous IEs, preferably from distant geographical regions, is important to demonstrate that identified anti-adhesion drugs inhibit the binding of IEs with diverse genetic background and are not specific to a single line only. Since the number of hits to test with IEs is low, the two-step procedure provides HT identification of anti-adhesion molecules.

DEVELOPMENT OF SCREENING PROCEDURE AND IDENTIFICATION OF NOVEL ANTI-ADHESION COMPOUNDS TO TREAT SM SYNDROMES CAUSED BY P. falciparum USING SMALL MOLECULE LIBRARIES

ChemBridge library. The development and proof-of-principle of our HT approach for the screening of anti-adhesion molecules have been performed [47] using a subset (n = 10,000) of the DIVERSet small molecule library from the ChemBridge Corporation. This library was selected based on its commercial availability, substantial size appropriate for identification of a sufficient number of active molecules, possibility to purchase separate subsets at affordable price for the initial development, as well as required physicochemical and pharmaceutical characteristics reflected in multiple publications. These characteristics include broad biologically relevant pharmacophore diversity space. For example, 50,000 drug-like small molecules of the DIVERSet library cover 60% of the total 3-point pharmacophore space of the entire EXPRESS-Pick™ Collection (internal) consisting of 500,000 compounds.

Before working with the small molecule libraries, we demonstrated the feasibility of our two-step approach using antibodies that inhibited the binding of three receptors, CD36 (the most common IE-binding receptor), CSA (receptor involved in PM), and ICAM-1 (receptor involved in CM) and erythrocytes infected with parasite lines binding to these receptors. When using PfEMP1 domains binding to these receptors and the corresponding binding parasite lines, we found a remarkable concordance between the in vitro molecular Bio-Plex assays.
and assays with live IEs, which strongly supported the feasibility of our two-step approach for screening anti-adhesion molecules. Since physical principles and readouts of the assay are the same for large (antibodies) and small anti-adhesion molecules, we expected to get similarly concordant results using the same two-step approach and small-molecule libraries.

As the first step, we used a sub-library of 10,000 compounds to prepare 125 pools, each containing 80 compounds (125 μM solution of each compound in DMSO) for the initial screening. In this way, the initial screening of all 10,000 compounds required less than two 96-well plates to assess their inhibitory effect on the binding of each of three endothelial receptors to the corresponding PfEMP1 domains.

We chose 75% inhibition of binding as an arbitrary threshold for our in vitro assay. Five different CD36-binding domains were selected for the screening of the CD36 binding inhibition. Interestingly, we did not find mixtures that inhibited CD36 binding over the selected threshold and, consequently, did not pursue further deconvolution. It is likely that identification of more efficient compounds for the inhibition of CD36 binding requires a larger library. For two other receptors (CSA and ICAM-1) and the corresponding three CSA-binding domains and one ICAM-1-binding domain (the only one identified by us in the NF54 line), we successfully identified inhibitory mixtures and deconvoluted them to three and two active individual compounds, respectively. Moreover, out of three compounds that inhibited the binding of the CSA-binding domain to CSA, two compounds shared the same sulfonyl-containing primary scaffold, which further strengthened the feasibility of our approach.

Using the in vitro assay, we demonstrated that the IC$_{50}$ values for both receptors were in a low micromolar range: 1.73 μM for the full-length VAR2CSA PfEMP1 protein that contained six CSA-binding domains [47], and 18 μM for the ICAM-1 binding domain PF11_0521 DBL2β3. The latter binds ICAM-1 receptor with a very high avidity (∼2 nM) [46, 59]. Therefore, identification of small molecular compounds that can compete in the domain–receptor interactions at low micromolar concentration is an important step toward identification of more potent compounds.

In accordance with our two-step approach, we demonstrated that compounds identified in our first in vitro step inhibited the binding of live IEs. In these experiments, which were performed at a single compound concentration of 100 μM as a proof-of-principle, we used the CS2 (CSA-binding) and 3G8 (ICAM-1-binding) parasite lines [74]. These lines have originated from a completely different genetic and geographic background (IT4 line from South-East Asia [75]) than the NF54 line (from Africa [76]) that was used as a source of our recombinant receptor-binding domains for the in vitro assays. Our experiments with live IEs clearly demonstrated that the identified active compounds inhibited the binding of CS2 and 3G8 lines to the corresponding binding receptors. This was a very important finding as it indicated that small molecule inhibitors may overcome a challenge presented by the domain diversity and can inhibit the binding of IEs of different genetic origin. Moreover, for the CS2 line, the identified compounds not only blocked, but also reversed the established interactions between IEs and CSA. This is also an extremely important feature, as reversing the established sequestration might have a significant impact on the SM pathology due to the removal of already sequestered parasites contributing significantly to the vascular inflammation.

Therefore, by using a subset (10,000 compounds) of the DIVERSet library, we not only successfully developed a two-step screening procedure, but identified three compounds that inhibited IE adhesion to the CSA receptor and two compounds that inhibited IE adhesion to the ICAM-1 receptor – two principal receptors involved in the sequestration of IEs in PM and CM [47]. This indicates that screening of currently commercially available full DIVERSet collection of small molecules from ChemBridge ($n = 150,000$) may reveal up to 30-50 individual active compounds/hits to select for the best leads based on their activity, cellular and animal toxicity, and pharmacokinetics/pharmacodynamics (PK/PD) in order to further develop them into anti-adhesion drugs.

**Natural product library (conopeptides).** Other libraries may provide an additional opportunity for screening for anti-adhesion compounds. In this respect, collections of natural products might be a good source for this type of screening. As a proof-of-principal, we used [77] a library of conotoxins/conopeptides, which are compounds expressed in the venom of marine mollusks belonging to the *Conus* genus. There are hundreds of thousands of these novel compounds, mostly disulfide-rich and well-structured peptides, expressed by more than 850 species of *Conus* mollusks [78-80]. Their presence in the venoms of mollusks does not necessarily mean that they all are toxic or harmful for humans. In addition, they can be modified to abolish the toxicity, while preserving their biological activity [81]. This class of molecules is actively pursued for the discovery of novel drugs to treat various diseases [82].

The high specificity and affinity of conotoxins for many ubiquitous somatic receptors suggest that they might be powerful compounds to prevent or reverse interactions between domains of malaria adhesins, PfEMP1 proteins, and host receptors involved in SM. In our work [77], we extracted venom samples from 460 specimens of *Conus nux* and fractionated them by size-exclusion (SE) high performance liquid chromatography (HPLC), followed by reverse phase (RP) HPLC. The obtained fractions were tested for the inhibition of binding between three vascular receptors (CD36, CSA, and ICAM-1) and the corresponding receptor-binding
PiEMP1 domains immobilized on BioPlex beads using the above-described in vitro approach [47]. Remarkably, six individual C. nux venom fractions, which contained either single peptides or a very limited set of peptides, affected the binding of several domains with different receptor specificity to the corresponding receptors, including proteins CD36 and ICAM-1 and polysaccharide CSA. These results suggest that either the peptides in these fractions bind to common structural elements in different PiEMP1 domains with a similar general scaffold (as we discussed above) or a few different peptides in the fraction interact with different domains (or receptors) and block the domain—receptor interactions. The ability of conotoxin fractions to inhibit the binding of IE adhesins with multiple binding specificity to a broad set of receptors can be extremely useful in the development of efficient and safe anti-adhesion drugs against SM [15, 16], because IEs isolated from SM patients often bind multiple receptors [57, 83]. Therefore, natural products, such as conopeptides, might be a promising source of anti-adhesion compounds that can be used for identification of leads for their development into anti-adhesion anti-malarial drugs.

TPIMS libraries. The above results demonstrate that screening of large libraries may provide a sufficient number of hits with low IC₅₀ values for their development into leads and, further, into anti-adhesion drugs; and that peptides and peptidomimetics might be extremely useful classes of molecules to identify these leads. Such large mixture-based libraries, ideally suitable for our two-step approach, have been developed at the Torrey Pines Institute for Molecular Studies (TPIMS) in a very convenient format allowing analysis of millions of compounds using just hundreds to thousands of samples [84–86]. The principle of this method is in the preparation of positional scanning libraries, each representing a different scaffold with several replacement groups, using the tea-bag approach [87]. The sub-libraries of each scaffold have one fixed (defined) replacement group and combinations of other replacement groups. The number of these sub-libraries is equal to the number of variants for the replacement groups. Analysis of screening results allows identification of the most active replacement groups. Then, a new sub-library(s) containing all combinations of these most active groups is prepared and tested for its activity. In 2–3 iterations (depending on the complexity of the initial library), a library of individual compounds is prepared based on the activity of mixtures with the defined replacement groups, and the most active individual compounds are identified. As we mentioned above, the screening of mixture-based libraries requires considerably less effort, time, and resources compared to the traditional HT screening of individual compounds.

TPIMS libraries only partially overlap with the structural space of currently approved drugs and, therefore, further add to the molecular complexity of libraries commonly used in HT screening [88, 89] and expand the traditional relevant space of medicinal chemistry [89]. In addition, large mixture-based libraries designed around a limited number of molecular scaffolds provide an opportunity for rapid elucidation of the structure—activity relationships (SARs) during screening. TPIMS collection is characterized by a larger degree of molecular complexity and larger 3D conformational space than typically available commercial collections [88] and possesses the drug-like properties similar to the property space of known drugs [89]. Successful application of TPIMS library collection in preclinical and clinical studies has been extensively reviewed [85, 90].

In our work [91], we screened a collection of TPIMS libraries consisting of more than 30 million compounds designed around 75 molecular scaffolds in order to identify the hits against binding between ICAM-1 (involved in CM) and the corresponding binding domain PF11_0521 DBL2β3 from the 3D7 line PiEMP1 protein [11, 46, 47], with IC₅₀ in a nanomolar range.

Based on the initial screening, dithiazole Positional Scanning Synthetic Combinatorial Library (TPI-2103, 3990 compounds) with the highest inhibitory activity and lowest complexity (two replacement groups) was selected for positional scanning deconvolution of the binding-inhibiting compounds. Individual compounds (104 in total) derived from the deconvolution of TPI-2103 represented all combinations of the selected most active R1 and R2 replacement groups. Their testing revealed two compounds (2648-40 and 2648-33) that inhibited ICAM-1 binding by more than 50% at 1 μM concentration. These two compounds were then characterized at a range of concentrations for the inhibition of ICAM-1 binding (i) in vitro using the bead-immobilized domain and (ii) with live erythrocytes infected by heterologous ICAM-1-binding 3G8 parasite line, as well as for cytotoxicity against two human cell lines.

The two compounds had the IC₅₀ values of 362 nM (2648-40) and 696 nM (2648-33) in vitro, which was about 50 and ~25 times, respectively, better than for the compounds identified in our previous work with the small-size ChemBridge sub-library [47]. Importantly, their inhibitory activity with live IEs was similar (IC₅₀ of 352 nM and 669 nM for 2648-40 and 2648-33, respectively). Thus, the same compound provided stronger inhibition of ICAM-1 binding on the beads and of binding between IEs and surface-immobilized ICAM-1, which was in accordance with our previous observations in [47]. Furthermore, the ICAM-1-binding domains of the PiEMP1 protein expressed on the surface of erythrocytes infected with NF54 line (used in in vitro experiments) and 3G8 line (used in experiments with live IEs) significantly differed in their amino acid sequence (44% identities, 55% positives, and 15% gaps by BLAST). Nevertheless, active compounds identified using a particular genetic background in the in vitro assays were successful in
inhibiting the binding of live IEs of very different genetic background, which again corroborated our results obtained using completely different compounds [47]. We believe that our approach might be universally employed for identification of anti-adhesion compounds against various receptor–domain pairs despite a substantial diversity of PiEMP1 domains/proteins.

The cytotoxicity studies demonstrated the selectivity index (SI, ratio of 50% cytotoxic concentration (CC50) toward human cell lines to 50% binding-inhibiting concentration (IC50) between the surface-immobilized receptors and live IE) of ∼70 and ∼37 for the two compounds, respectively. This is an excellent result, as SI > 10 is considered the minimally acceptable value and SI > 50 is an ideal value for the target candidate profile (TCP-1) [48], because after getting into the bloodstream, common malaria parasite-killing drugs should cross at least 3 membranes (red blood cell membrane, parasitophorous vacuole membrane, and parasite plasma membrane) for their effect. The anti-adhesion drugs are supposed to work immediately at the outer surface of IEs after entering the blood. Therefore, the two identified lead compounds provide an excellent starting point for the development of anti-adhesion drugs against SM and CM, in which IE adhesion to ICAM-1 receptor plays an extremely important pathological role.

As we mentioned above, the binding to ICAM-1 and EPCR is implicated in the development of SM and CM. Many parasites isolated from SM or CM patients demonstrated the presence of a tandem of ICAM-1- and EPCR-binding domains in the expressed PiEMP1 molecules [59, 60, 92-96]. Even if both these domains in a tandem are required for the development of SM and/or CM, this does not mean that the drugs should inhibit the binding of both domains to the corresponding receptors to successfully block IE sequestration. In this case, inhibition of ICAM-1 binding only may significantly reduce the efficacy of IE sequestration, especially in the brain microvasculature, where EPCR is expressed at low levels, while the content of ICAM-1 is high and further increases during brain inflammation (reviewed in [97]), in particular, during CM-associated inflammation induced by TNFα (reviewed in [98]). In this respect, identification of other compounds that will not only prevent but also reverse ICAM-1 sequestration, may have a significant advantage in the anti-adhesion adjunct treatment. As demonstrated in our studies and works of other authors, ICAM-1 binds to the ICAM-1-binding DBL domains with an extremely high avidity (2-4 nM) [46, 59]. Therefore, identification of such compounds might not be a trivial task. Most likely, they will have an allosteric, and not competitive mode of action. The leads identified in [47, 91] did not reverse the established ICAM-1–domain interactions. Hence, molecules with this activity are still to be discovered. Nevertheless, just inhibition of ICAM-1 binding might be sufficient to reduce or prevent the development of pathologies related to the continuous IE sequestration in SM or CM, although this hypothesis should be clarified experimentally.

CHALLENGES IN TESTING ANTI-MALARIAL ANTI-ADHESION DRUGS IN PRE-CLINICAL STUDIES

Since anti-adhesion drugs do not usually kill malaria parasites (if they did, this would be an additional important bonus), it is difficult to justify their trials in human patients with severe forms of malaria. One of the effects these drugs should produce is reduction of inflammation of endothelial cells (ECs) by preventing or reversing IE cytoadhesion. This can be studied in vitro by incubation of ECs with IEs that can adhere to these cells [for example, CSA-binding IEs and human trophoblast-derived BeWo cells expressing CSA, or ICAM-1 binding IEs and TNFα-stimulated human umbilical vascular endothelial cells (HUVEC) that express ICAM-1] in the presence and absence of anti-adhesion compounds, followed by detection of inflammatory molecules secreted by the cells. However, these effects are not easy to demonstrate in vivo in the preclinical studies, as no appropriate animal model for P. falciparum malaria is available. PiEMP1 proteins are specific for P. falciparum only, while malaria parasite species that infect rodents are very different and do not share homologous PiEMP1 proteins. Even humanized mouse malaria model [99], in which immunodeficient laboratory NOD-scid gamma (NSG) mice are injected with human erythrocytes for several days to substitute mouse erythrocytes and then challenged with P. falciparum to establish the infection, will not be appropriate, as mouse endothelial receptors, though homologous, are different from the human receptors interacting with PiEMP1 proteins and may not provide adequate IE cytoadhesion. Therefore, appropriate models should be designed for testing anti-adhesion compounds (beyond their cellular, animal, and human cytotoxicity) before moving to trials in humans. Potentially, some non-human primate models, like P. falciparum infection of Aotus monkeys [100], may be used to test IE sequestration and activity of anti-adhesion drugs.

IMPLICATIONS OF ANTI-ADHESION DRUGS FOR OTHER TARGETS

The two-step approach developed by us for the HT identification of anti-adhesion compounds might be useful in the identification of compounds for treatment of other diseases, for which interactions between two known molecules/molecular structures play an important role in the disease pathology, e.g., infection of cells by viral, parasitic, or bacterial pathogens through specific receptors.
Identification of anti-adhesion compounds from large libraries via HT screening in vitro followed by their in vivo or ex vivo testing may significantly speed up the discovery of novel therapeutics for various diseases.

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