Development of \textit{Plasmodium falciparum} Protease Inhibitors in the Past Decade (2002–2012)

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Abstract: New drug targets for the development of antimalarial drugs have emerged after the unveiling of the \textit{Plasmodium falciparum} genome in 2002. Potential antimalarial drug targets can be broadly classified into three categories according to their function in the parasite’s life cycle: (i) biosynthesis, (ii) membrane transport and signaling, and (iii) hemoglobin catabolism. The latter plays a key role, as inhibition of hemoglobin degradation impairs maturation of blood-stage malaria parasites, ultimately leading to remission or even cure of the most severe stage of the infection. Intraerythrocytic \textit{Plasmodia} parasites have limited capacity to biosynthesize amino acids which are vital for their growth. Therefore, the parasites obtain those essential amino acids via degradation of host cell hemoglobin, making this a crucial process for parasite survival. Several plasmodial proteases are involved in hemoglobin catabolism, among which plasmepsins and falcipains are well-known examples. Hence, development of \textit{P. falciparum} protease inhibitors is a promising approach to antimalarial chemotherapy, as highlighted by the present review which is focused on the Medicinal Chemistry research effort recorded in the past decade in this particular field.

Keywords: \textit{Plasmodium falciparum}, hemoglobin catabolism, plasmepsin, falcipain, falcilysin, aminopeptidase, proteases, antimalarial.

1. INTRODUCTION

Malaria is one of the most threatening infectious diseases that mainly affects the world’s poorest countries in tropical areas [1]. Approximately 40% of the world population lives at risk of this disease and this constitutes a large burden on the health and economic development of low-income countries [1-3]. Although the World Malaria Report 2011 showed that there has been significant and durable progress in battling this disease, emergence of parasite resistance to antimalarial medicines remains a threat to this continued stride toward the reduction of malaria cases in the world. There are five species of \textit{Plasmodium} which cause malaria in human. Those species are \textit{P. ovale}, \textit{P. malariae}, \textit{P. knowlesi} [4, 5], \textit{P. vivax} and \textit{P. falciparum (Pf)}, the latter being the deadliest [6, 7]. Malaria is transmitted by a female infected \textit{Anopheles} mosquito that harbors \textit{Pf} sporozoites in its salivary glands; by biting a human, the mosquito injects those sporozoites into the blood-stream, and they are then carried to the liver to develop into merozoites, which subsequently are released into the blood stream invading host red blood cells and multiplying asexually as trophozoites. Some erythrocytic parasites develop into male and female gametocytes, the only forms able to infect other mosquitoes and there reproduce sexually, permitting completion of the life cycle [2].

Currently, there are drugs targeting different stages in the malaria life cycle such as chloroquine 1, artemisinin 2, and primaquine 3 (Fig. 1). However, the increasing spread of parasite strains resistant to currently used antimalaria has put enormous pressure on public health systems to introduce new treatments [8]. Given that the hope for a long-lasting vaccine against malaria is yet an unmet goal, it appears that control of the disease has to rely mostly on chemotherapy in the foreseeable future [9].

![Fig. (1). Structure of known antimalaria: chloroquine (1), artemisinin (2), and primaquine (3).](image-url)

Since the unveiling of the \textit{Pf} genome, a decade ago, new targets for development of antimalarial drugs have arisen [10]. Antimalarial targets can be broadly classified into three
categories according to their function in the parasite’s life cycle: (i) biosynthesis, (ii) membrane transport and signaling, and (iii) hemoglobin catabolism. To the first target category belong Pf enzymes in charge of generating nutrients required for malaria parasite growth. For instance, there are different biosynthesis pathways today used as targets for the discovery of antimalarial drugs such as the folate biosynthesis pathway [11-13]. The second target category includes pathways which mediate the uptake of nutrients into cells and the generation and maintenance of transmembrane electrochemical gradients, for instance, the plasmoidal surface anion channel (PSAC) [14-16]. Other examples which fall within this target category are enzymes whose substrates are involved in intracellular signal transduction, for instance, farnesyltransferase [17-19]. Finally, targets within the third target category play an essential role in the development of intraerythrocytic malaria parasites. Since the digestion of hemoglobin is presumably an essential catabolic function performed by the blood stage parasites, the proteases participating in this pathway have been proposed as targets for the development of novel antimalarial drugs [20-22]. This review will mainly focus on such proteases and their inhibitors developed in the past decade.

2. HEMOGLOBIN CATABOLISM

During the intraerythrocytic development of the asexual stages of Pf, the parasite inhabits a parasitophorous vacuole (PV) formed inside the red blood cell (RBC); host cell hemoglobin is endocytosed by the PV and transported to an acidic compartment known as the parasite’s food vacuole (FV). Within the FV, proteases degrade most of the host hemoglobin to free amino acids that can then be incorporated into newly synthesized proteins [23] and used to modulate the osmotic status of the host cell [24, 25]. In addition, removal of hemoglobin also frees up space within the RBC for parasite replication [26]. Early in the degradative pathway, free heme is released and oxidized from the ferrous (Fe^{2+}) state to the ferric (Fe^{3+}) hematin. Both heme and hematin are potentially toxic to the parasite [27]. To counter this, the parasite has evolved a detoxification system resulting in the formation of the hemozoin pigment, an inert crystalline β-hematin polymer [28, 29]. Antimalarial 4-amino-quinolines, such as chloroquine (I, Fig. 1), appear to function by disrupting this sequestration, leading to an accumulation of toxic heme products [30].

Studies suggest that the proteases in charge of hemoglobin degradation are essential for the parasite growth since the parasite has limited ability for de novo biosynthesis of the amino acids needed for its own proteins. It has been shown that Pf can also obtain amino acids exogenously, as parasites can survive in media supplemented only with isoleucine, an amino acid not contained in hemoglobin; yet, in the absence of an exogenous amino acids source, the parasite can solely rely on hemoglobin degradation to obtain most of the required amino acids [31]. Moreover, it has been clearly shown that Pf utilizes hemoglobin as an amino acid source for protein synthesis, as amino acids from hemoglobin degradation have been detected in parasite’s proteins, and in vitro experiments where hemoglobin digestion has been impaired revealed that parasite development and morphology were significantly affected [32].

Extensive study of this protease cascade has resulted in the model outlined in (Fig. 2) for hemoglobin degradation. There is evidence that the degradative enzymes function in a semi-ordered pathway [30], with aspartyl proteases, being the first to participate in this proteolytic pathway by making the initial cleavage in intact hemoglobin. Subsequent degradations of globin into small peptides chains are carried by cysteine proteases, metalloprotease falcilysin (FLN), and dipeptidyl amino peptidase 1 (DPAP1) [33]. Following, the small peptides are transported to the parasite cytoplasm where they are terminally degraded to amino acids by exopeptidases (Fig. 2) [33].

From the above, it is understandable that proteases involved in this catabolic pathway have become of substantial interest for the malaria research community. Certain inhibitors for these enzymes have been tested in in vitro experiments and found to block Pf growth and its expansion to other erythrocytes [35-37]. Therefore, the search for potent inhibitors of these proteases has become a new strategy for developing new antimalarial drugs.

Fig. (2). Proteolytic cascade in hemoglobin degradation by Pf parasites. For a more detailed explanation refer to: http://primweb.cc.huji.ac.il/malaria/FrameHemoglobindeg.html and see ref [34].

2.1. Aspartyl Proteases (Plasmepsins)

Pf contains at least 10 aspartyl proteases, known as plasmepsins: plasmepsins I, II, and IV–X, and histidine aspartic protease (HAP). The precise role of each of the plasmepsins in the parasite metabolism is not clear. To date, the most extensively studied plasmepsins are plasmepsin I (PIm I; EC number: 3.4.23.38), plasmepsin II (PIm II; EC number: 3.4.23.39), plasmepsin IV (PIm IV; EC number: 3.4.23.B14) and HAP. These aspartyl proteases are expressed during the erythrocytic stage of the parasite [38], have all been shown to be directly involved in the process of hemoglobin degradation [38, 39], and have all been characterized structurally [40]. High levels of sequence homology (60%-70%) [41] are observed between PIm I, II, IV, and HAP, which also lies in a cluster on the same gene [39]. Compared to PIm II, the binding site regions of PIm I, IV, and HAP show 84%, 68%, and 39% identity, respectively [42].
Plm I and Plm II catalyze the initial step in the breakdown of hemoglobin by the parasite by making the first cleavage of hemoglobin between Phe33 and Leu34 of the α-chain to generate globin and free heme [38, 39]. In addition to its participation in hemoglobin degradation, the interest in Plm IV is also motivated by the fact that it is the only plasmepsin located in the FV of Pf which has orthologs in the other Plasmodium species infecting human [41]. HAP is unique in the sense that it has a histidine in the place of the first canonical aspartic acid. Whether this results in an aspartic- or serine-protease-like mechanism has been subject of discussion [43]. Computational predictions indicate that only the aspartic acid (Asp214) is directly involved in catalysis, while the histidine residue (His34) provides critical stabilization along the cleavage [44]. Recent knock-out studies revealed that single, double, and even triple knock-outs of FV plasmepsins are viable, suggesting that multiple plasmepsins must be targeted to produce antiparasitic effects [45]. This also indicates that there is a high number and functional redundancy of plasmepsins in the hemoglobin pathway. However, it has been shown that Plm inhibitors significantly attenuated parasitemia both in culture and in animal models [46].

As the crystal structure of Plm II was the first one to be publicly available, most research focused on the development of Plm II inhibitors. Plm II contains 329 amino acids and its binding cleft, also known as the catalytic dyad, is constituted by Asp34 and Asp214, which are bridged by a water molecule [47]. Hydrolysis of the peptide bond between hemoglobin’s Phe33 and Leu34 takes place, and a water molecule [47]. Hydrolysis of the peptide bond between hemoglobin’s Phe33 and Leu34 takes place once Asp214 abstracts one of the protons from the water molecule, and generates two peptide chains (Fig. 3) [35, 47]. The cleavage mechanism concludes when the peptide products leave the active site and the water bridge between Asp34 and Asp214 is regenerated.

Plm I and II have a 73% sequence homology suggesting that both enzymes can be inhibited by related molecules. Therefore, Plm I and II have become the main targets of most plasmepsin inhibitors developed as potential antimalarial drugs. Yet, these plasmepsins show high structural similarity with human cathepsin D, which makes selectivity an important factor to take into account when designing suitable inhibitors.

Well known HIV-1 aspartyl protease inhibitors, such as ritonavir, indinavir, nelfinavir, saquinavir, atazanavir and amprnavir, were also found to inhibit Pf aspartyl proteases and the development of parasites at pharmacologically relevant concentrations [48]. All of these inhibitors undergo non-covalent interactions with the proteases and share key structural features for the inhibition of aspartyl proteases, which is a hydroxyl or hydroxyl-like moiety that coordinates to the catalytic dyad and mimics the transition state for peptide bond hydrolysis. The (S)-hydroxyl group displaces the water molecule from the catalytic site and forms a hydrogen bond with Asp [49, 50]. Like these HIV-1 aspartyl protease inhibitors, most Plm I and II inhibitors mimic the tetrahedral intermediate formed during the aspartyl protease catalysis. There are several transition state analogue cores used for the design of Plm inhibitors [35, 51-65], but the most important include the statine core [54, 56, 57, 64], the reversed-statine core [53, 54], or a hydroxyethylamine motif (Fig. 4) [56, 61, 62].

Studies using encoded combinatory libraries based on the statine core structure 4 (Fig. 5) allowed determining that β-branched side chains are preferred in P2 and hydrophobic side chains as phenyl or isobutyryl in P1. In addition, P2 and P3 substituents impart selectivity in Plm II inhibitors (Fig. 5) [66, 67]. One advantage of this type of inhibitors is that they don’t cross inhibit other proteases such as serine-, cysteine-, or metallo-proteases [68].

Recently reported statine-based inhibitors include the one found by Bosisio and co-workers [64]. They coupled a series of statine-based inhibitors with primaquine 3 and found low nanomolar inhibitors of Plm II with IC50 between 0.59 to 400 nM and low micromolar activity in vitro against Pf, the best of which (5) is shown in (Fig. 6). A direct correlation between the compounds’ activity against Plm II and the in vitro parasite growth suggested that the main mechanism of these inhibitors was Plm II inhibition and consequently, the digestion of hemoglobin that is essential for Pf survival as stated before. Compound with the linker derived from a succinic acid was the least active of the series which is in agreement with the finding suggesting that an aromatic substituent is preferred in P3 for Plm II inhibition [66]. The introduction of an aromatic ring as a linker in general structure 6 increased the activity as expected and compounds with a naphthyl were more active against Plm II. Still, there was not significant increase of the resulting derivatives against parasite growth. Compound 5 was considerably more active against the enzyme and parasite growth with IC50 0.59 nM on in vitro in-

Fig. (3). Schematic mechanism of hemoglobin cleavage by Plm II.
hibition of Plm II, and IC$_{50}$ of 0.4 µM on in vitro inhibition of growth of PfD10 [64]. Additional studies were done to verify the dual-action of compound 5. The products of the hydrolis of Lys-Leu peptide bond in compound 5 were assessed as inhibitors of the parasite development and Plm II activity. Even though such products were both active against parasite growth in the micromolar range, compound’s activity against the parasite development was found to be mainly due to Plm II inhibition.

Samuelsson and co-workers reported [54] a series of statine-reversed core inhibitors of Plm I and Plm II. The best inhibitor 7 of the series (Fig. 7) exhibits $K_{i}$ values of 250 nM and 1.4 µM for Plm I and II, respectively. To increase affinity to both plasmepsins a benzyl group was added to the N-terminus to interact with the S1’ pocket of the proteases. Furthermore, a hydrazine moiety was inserted to allow flexibility to the benzyl substituent in the N-terminus to better fit in the S1’ pocket. All the compounds were tested against Plm I and II activity. From the inhibition results it can be inferred that the N-benzyl derivatives were almost inactive whereas the aza-benzyl derivatives promote the inhibition of both plasmepsins. Also, the presence of the carboxybenzyl group in the P2’ position seems to increase the activity of the compounds. Samuelsson and co-workers [54] also reported that the basic piperidine- and pyridine-substituents are present in the majority of the potent inhibitors, for instance, see compound 7.

In 2003, Nöteberg and co-workers [61] reported a series of compounds, including a basic hydroxyethylamine transition state isostere, designed and synthesized as inhibitors of Plm I and Plm II. These compounds were designed on the basis of previous findings from Ellman’s group, who determined that large substituents were suitable for P1’ position [69]. In comparison to Ellman’s compounds, Nöteberg’s had the nature of the peptidic inhibitor minimized using a single prime side amino acid residue bearing a biphenyl side chain. This yielded compounds which were highly selective for plasmepsins over cathepsin D, the most active of which (8) is displayed in (Fig. 8).

In the same year, the same group also synthesized and screened compound libraries based on the general structure 9, and found inhibitors with $K_{i}$ values in low nanomolar range that targeted the malaria proteases Plm I and Plm II, again with high selectivity versus cathepsin D (Fig. 9) [62]. This study allowed not only to find very active and selective inhibitors, like 10 (Fig. 9), as well as to demonstrate that

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**Fig. (4).** Transition-state mimicking groups in peptidomimetic plasmepsin inhibitors: reduced amine [52, 59], statine [54, 56, 57, 64], hydroxypropylamine [51], reversed-statine [53, 54], dihydroxyethylene (C- and N-duplicated) [35, 55, 63], norstatine [58, 60, 65], and hydroxyethylamine [56, 61, 62].

**Fig. (5).** General structure requirements for Plm I and Plm II inhibitors based on the statine core.
very diverse side chains in P1’ and P3 positions are suitable for Plm I and II inhibition.

Fig. (7). The most active Plm I and Plm II inhibitor, based on the statine-reversed core, designed by Samuelsson and co-workers [54]. Bn=Benzyl; Cbz=Benzyloxycarbonyl.

Fig. (8). The most active Plm I and Plm II inhibitor of the biphenyl series developed by Nöteberg and co-workers: \( K_i(\text{Plm I}) = 115 \text{ nM} \); \( K_i(\text{Plm II}) = 121 \text{ nM} \) [61].

Most of the reports on Plm II inhibitors describe non covalent interactions with the aspartyl proteases. However, there are a few examples of irreversible inhibitors in the literature [70-72]. Woster and co-workers [71] studied the synthesis and screening of Plm II inhibitors and found that three compounds produced irreversible inactivation of the enzyme with IC\(_{50}\) values in the low nanomolar range. They hypothesized that an (5)-hydroxyl substituent moiety bearing a latent electrophile should act as an irreversible inhibitor of Plm II. For instance, Asp214 or an adjacent basic amino acid could abstract an acidic proton from the inactive electrophile resulting in the generation of an \( \alpha,\beta \)-unsaturated system (11), ketenimine (12), or an allene (13 and 14) in the catalytic site (Fig. 10).

Fig. (9). Core structure 9 of the library of potential Plm I and Plm II inhibitors studied by Nöteberg’s team. Compound 10 was the most active and selective inhibitor, with \( K_i(\text{Plm I}) = 12 \text{ nM} \); \( K_i(\text{Plm II}) = 110 \text{ nM} \); and \( K_i(\text{Cathepsin D}) = 3300 \text{ nM} \) [62].

Fig. (10). Structures of irreversible Plm II studied by Woster’s group [71].
Woster’s group found compounds 11, 13, and 14 to be active in nanomolar range between 20-350 nM. Interestingly, compound 14 showed the greatest selectivity for Plm II over cathepsin D despite the fact that the latent electrophile was not adjacent to the (S)-hydroxyl moiety. Compound 11 was a poor inhibitor of the parasite activity in vitro in the blood stage, contrary to compounds 13 and 14 which displayed IC\textsubscript{50} values of 7.7 and 9.2 μM, respectively, in the infected erythrocyte assay. Still, the mechanism through which compounds 11, 13, and 14 exert their activity against the enzyme is unknown. Thus, additional studies are required in order to explain the specific amino acid residues which participate in the covalent bond formation.

In general, the aforementioned peptidomimetic inhibitors exhibit low nanomolar activity in vitro against plasmepsins, however their in vitro activity against blood stage parasites usually drops significantly. In other words, effective inhibition of plasmepsins was found to correlate poorly with parasite killing in culture [73]. This frequent effect might be attributed either to compounds possessing unfavorable pharmacokinetic properties, e.g. poor cell permeability, or to the fact that not all four plasmepsins are inhibited equally well.

Although most Plm I and II inhibitors are peptidomimetic compounds that mimic the tetrahydridal intermediate formed during the aspartyl protease catalysis, some nonpeptidic molecules have generally been more successful than the peptidomimetic inhibitors in demonstrating high activities in cell culture [74]. This may be the result of more appropriate structural profiles for membrane permeability. A number of nonpeptide plasmepsin inhibitors displaying low nanomolar range against parasite growth in vitro (Fig. 11) have been reported [75].

Via a high-throughput screening of the Roche compound library, substituted piperidines were identified and further developed as inhibitors of the aspartyl protease rexin [76]. Subsequent structural optimization of the initial lead structure, found to inhibit Plm I and Plm II with an IC\textsubscript{50} value of about 1 μM for both variants, resulted in compounds, such as compound 15 (Fig. 11), exhibiting nanomolar activity in cellular assays [77].

Another study, reported by Boss and co-workers [78], of high-throughput screening of a commercial library using fluorescence resonance energy transfer (FRET) assays to measure the Plm II activity, identified low μM inhibitors such as type I and type II (Fig. 12) as promising compound classes. Further optimization of the compounds was able to increase the activity of type I and type II inhibitors by 250-fold and a factor of 60, respectively. Such improvements led to compound 16 (Fig. 11) and 17 (Fig. 13), respectively [77, 78]. The binding modes of the 4-aminopiperidines (compound 16) were elucidated by X-ray crystallography (PDB-code: 2BU) [79], which certainly will facilitate the subsequent structure-based drug design of this promising class of compounds.

For type II inhibitors, replacement of the n-pentyl chain by shorter ones, as well as the insertion of a heteroatom in the chain, leads to a decrease on the activity of the inhibitors against Plm II. Still, little is known about the consequences on the activity of type II inhibitors upon alterations of R\textsuperscript{3} and R\textsuperscript{4} substituents. Boss and co-workers [78] demonstrated that hydrophilicity and polarity in the bis-heteroaryl moiety could be added to improved activity against the parasite growth. Their studies also showed that the aryl-amide unit and the aryl-amine are beneficial for global activity. Compound 17, an aryl-amine derivative shown in (Fig. 13), was found to be one of the most promising inhibitors against Plm II and parasite growth of the series of compounds considered, having IC\textsubscript{50} values of 374 nM against Plm II and 273 nM against the parasite development.

In addition, other non peptidic inhibitors of plasmepsins include the achiral oligoamines reported by Diederich and co-workers [80]. They developed an initial library of 11 compounds and tested them against six different aspartyl proteases. Several hits were identified and some of them were found to be also selective. They demonstrated that the sulfonamides acceptor groups are preferred over the carbonyl moiety because they allow additional degrees of freedom. Polymethylene links confer flexibility to the oligoamine derivatives, which requires a high degree of pre-organization from the molecules before binding to the protein. One of the most active inhibitors of this series against Plm II, compound 18 shown in (Fig. 14), exhibited K\textsubscript{i} value of 7.0 μM.

Noteworthy, several of the peptidomimetic and nonpeptide plasmepsin inhibitors have been assessed against both chloroquine-resistant and chloroquine-sensitive strains of P\textsubscript{f} with similar potencies against both strains, indicating no cross-resistance between chloroquine and plasmepsin inhibitors [64, 75].

2.2. Cysteine Proteases (Falcipains)

After aspartyl proteases carry out the first cleavage of the hemoglobin into two peptide chains, falcipains (FPs) come to

![Fig. (11). Non-peptidic Plm II inhibitors with P/IC\textsubscript{50} values of 50 nM (15) and 252 nM (16) [75, 77].](image-url)
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play their role to transform those peptides into smaller ones. FPs are papain-like cysteine proteases and the best characterized Cys proteases of the malaria parasite, in which a Cys catalyzes protein hydrolysis via nucleophilic attack to the carbonyl carbon of a susceptible bond (Fig. 15). Analysis of the Pf genome sequence suggested the existence of four FPs: FP1, FP2 (EC number: 3.4.22.B69), FP2', and FP3 [81].

Fig. (12). General structure of type I and type tertiary amines as inhibitors of Plm II [78].

Fig. (13). Compound 17, one of the most active aryl-amine derivatives against Plm II and parasite development with IC\textsubscript{50} values of 374 nM and 273 nM, respectively [78].

Fig. (14). Compound 18, one of the most active oligoamine derivatives against Plm II with \(K_c\) value of 7 \(\mu\)M, reported by Diederich and co-workers [80].

FP1 is distantly related to the other FPs in terms of sequence (<40% amino acid identity) and its exact physiological role has yet to be elucidated. Some studies suggested that FP1 could be important in oocyst production during parasite development in the mosquito midgut [82] and also could help in the invasion of the host cell by Pf. However, disruption of the FP1 gene does not disturb the parasite growth in the blood stage of the malaria life cycle [83]. Furthermore, a study showed that FP1 knockout parasites developed normally in erythrocytes, suggesting that this protease alone is neither required for parasite invasion nor for intracellular development within erythrocytes [84]. FP2, FP2' and FP3 are closely related and appear to be the key hemoglobinases in the acidic FV. FP2 is the most studied and FP2' is thought to arise by gene duplication, presenting 93% a.a. similarity with FP2 [85].

Fig. (15). Mechanism of peptide cleavage mediated by falcipains, Cys proteases of Pf.

FP2 and FP3 are single polypeptide chains presenting a high similarity in sequence (68% identity) and sharing similar sized prodomains. Their catalytic site has Cys and His residues (Fig. 16), whose side chains form a thiolate/imidazolium ion pair, and has also an Asn, responsible for correct orientation of the ion pair [81]. FP2 and FP3 contribute equally to hemoglobin degradation and both require reducing environment and acidic pH for optimal activity. Interestingly, the disruption of FP2 gene revealed that the loss of this enzyme alone is not sufficient to cause parasite lethality, thus suggesting the participation of additional cysteine proteases for parasite invasion and growth in human erythrocytes [86]. In contrast, the FP3 gene could not be disrupted, revealing that FP3 is essential to erythrocytic parasites and therefore indicating that efforts to develop cysteine protease inhibitors as antimalarial drugs should probably be focused on FP2 and FP3 [86]. The structural similarities of FP2 and FP3 can be a reason for these proteases performing similar functions in the FV. One difference between FP2 and FP3 is that FP3 showed to be far less active than FP2 against peptide substrates. However, FP3 is more active and stable at acid pH (as in the parasite’s FV) than FP2 [87]. Also, it has been estimated that the concentration of FP2 in trophozoites is 1.8 times that of FP3. Still, FP3 processes globin two times faster than FP2 does [88]. The structures of FP2 and FP3 forming complexes with known inhibitors of other Cys proteases have been reported: the FP2 crystal structure was reported in a complex with inhibitor E64 19 (PDB code=3BPF) [83] and the crystal structure of FP3 in a complex with Leupeptin 20 (PDB code=3BPM) [83] (Fig. 16) and with a vinyl sulfone (PDB code=3BWK) [89]. Both FP2
and FP3 showed to have a preference for substrates with a hydrophobic residue, especially leucine, at P2 position.

Although both FP2 and FP3 contribute almost equally to the digestion of hemoglobin, FP3 is usually less amenable to inhibition by peptidyl-based small molecules [83]; consequently, there is a variety of FP2 inhibitors in the literature [86]. In general, FP inhibitors can be classified in peptidyl-based inhibitors (covalent irreversible and reversible inhibitors) [83, 90, 91], including peptidomimetic compounds [92, 93], and nonpeptidic small inhibitors [94-96].

**Fig. (16).** Structures of known Cys protease inhibitors E64 (19) and Leupeptin (20).

### 2.2.1. Peptidyl-Based Inhibitors

#### 2.2.1.1. Irreversible Inhibitors

Most of the falcipain inhibitors identified so far are peptidyl-based molecules [86]. Given that Cys proteases have their enzymatic roles dependent on a catalytic Cys residue, an electrophilic warhead is necessary to inhibit these proteases. Among such warheads, prone to bind in an irreversible way to FPs, like fluoromethyl ketones [97], epoxysuccinyl [98] or azirine [98] derivatives, the most used are Michael acceptors that irreversibly inactivate Cys proteases via alkylation of the side chain thiolate of the Cys residue (Fig. 17) [37]. These Cys protease inhibitors include an α,β-unsaturated conjugated system such as vinyl sulfone or α,β-unsaturated carbonyls [99, 100].

**Fig. (17).** Schematic view of irreversible alkylation of a Cys via Michael addition.

Since Rosenthal and co-workers [100] reported in 1996 that vinyl sulfone inhibitors of FP2 blocked the development of PF1 in culture and presented antimalarial activity in vivo, there have been several efforts to synthesize Cys protease inhibitors. The best known attempt was done by the same research team in 2003 [101], when they reported the structure-activity relationships (SAR) for a series of peptidyl vinyl sulfones, vinyl sulfone esters, and vinyl sulfonamides as inhibitors of FP2 and FP3. In that study, they demonstrated that the SAR for the two proteases is similar and found multiple compounds to be potent inhibitors of both falcipains, especially amongst the vinyl sulfones (21, Fig. 18).

Studies of Rosenthal’s group showed that the SAR of FP2 and FP3 are similar and several compounds are potent inhibitors of falcipains and parasite growth. Also, the SAR for the inhibition of the proteases differed notably for those of inhibition of parasite development. In the SAR regarding the proteases, Rosenthal’s team found that inhibitors with the dipeptide core Leucine-homoPhenylalanine (Leu-hPhe) in P2-P1 provided potent inhibition for FP2, FP3 and for development of malaria parasites in vitro, and that potency was imparted by alterations of amino (P3)- and carboxy (P1’)-terminal substituents of vinyl sulfone. Still, substrates with Leu at P2 showed to be less active against FP3 than against FP2. Compounds with the core sequence Phe-hPhe exhibited modest FP2 inhibitory activity and changes at position P3 in these molecules had relatively low impact on activity. Compounds with the dipeptide sequence Phe-O-(phenyl)Ser had inhibitory activities with IC50 against FP2 from 56 to 290 nM. The same group then tested 30 compounds with the core sequence Phe-hPhe, which differed at the P1’ and P3 position. Vinyl sulfonamides and vinyl sulfones were seen to be less active against FP2 than the vinyl sulfonate esters. Still, inhibitory activity against parasite growth was higher for vinyl sulfones. Finally, these authors also found a pattern of activity against the parasite cultures, depending on the substitution in the aryl ring of the vinyl sulfonate esters: presence of the electron-donating methoxy group led to an increase of activity over the unsubstituted ring, whereas the last one showed to be more active than the one bearing an electron-withdrawing fluoride [101]. More recently, a computational 3D-QSAR study on these three vinyl sulfone, sulfonamide and sulfonate ester families has identified critical regions where any change in the steric, electrostatic, and hydrophobic fields of the molecules may affect the inhibitory activity [102]. For instance, bulky groups at R2 position (Fig. 18) tend to decrease biological activity while electropositive groups at R2 backbone, R1 side-chain and near the SO2–R1’ group are preferred to improve FP2 inhibitory activity. Also, compounds that orient any hydrophobic part of R3 group towards Tyr78 and Leu84 would exhibit enhanced FP2 inhibitory activity. Additionally, they also postulated that the general rank order of FP2 inhibitory activity, sulfonate esters > sulfonamide > sulfones, observed when comparing compounds that only differ on R1’ substituent could be related to the flexibility of this substituent.

Though peptidyl-based inhibitors like those above inhibit enzymatic activity of FPs at very low nanomolar range, their utility as therapeutic agents may be limited for their susceptibility to protease degradation and their poor absorption through cell membranes, so peptidomimetic scaffolds have been developed to bypass this problem [86]. A common strategy to avoid the therapeutic limitations of peptide-based
Fig. (18). Structure of vinyl sulfone inhibitors of FP2/3 with nanomolar activity against intra-erythrocytic P/ parasites [101].

inhibitors is to lock a defined conformation of the peptide into a rigid scaffold. There are several examples of peptidomimetic inhibitors of FP2 reported in the literature [92, 93, 103-106]. O’Neill and co-workers designed and synthesized novel 2-pyridone peptidomimetic FP2/3 inhibitors with chemical structures shown in (Fig. 19) [107]. All of these molecules displayed antimalarial activity in the micromolar range, being 22 the most active inhibitor of the series with an IC50 (3D %) value of 5.7 μM, followed by the α,β-unsaturated methyl ester 23 with an IC50 (3D %) value of 27.9 μM. Compound 24 was less active than their counterparts bearing the unnatural segment of hPhe residue. However, only 24 expressed activity against FP2/3, whereas the low solubility of 22 prevented its evaluation as a potential FP inhibitor.

Fig. (19). FP2/3 peptidomimetic inhibitors studied by O’Neill and co-workers [107]. Cbz=Benzylloxycarbonyl.

Other peptidomimetics bearing Michael-acceptor moieties for irreversable FP inhibition have been explored as antimalarials. For instance, Moreira and co-workers have developed vinyl sulfamides [108], aza vinyl sulfones [109], squaric acid [110] and 3-methylene substituted indolinone derivatives [111]; only some of the latter compounds were able to inhibit P/parasites in vitro at IC50 as low as 140 nM, the remaining presenting IC50 values in the micromolar range. In general, none of these compounds was able to exhibit potent FP inhibition activity.

Peptidomimetic nitriles have also been reported in the literature as potential FP inhibitors. Nitriles are known to inhibit cysteine proteases by the formation of a covalent thioimidate adduct resulting from the nucleophilic attack of the active site Cys residue [112]. Using a rational structure-based molecular modeling focusing on optimal occupancy of the main apolar FP2 pockets to obtain potent and particularly selective inhibitors, Diederich and co-workers [103] were able to design nitrile inhibitors of FP2 action, and showed that the ideal occupation of the selectivity-determining S2 pocket and the balanced electrophilicity of the nitrile group seem to be essential to achieve activity and selectivity. The 3D modeling studies specifically revealed that 1-aminocyclohexanecarboxylic acid should be the central core of the nitrile derivative to address the active site of FP2. The cyclohexyl motif should orientate the other substituents in the molecules to promote their interactions with the S2 and S3 pockets. They synthesized the nitriles derivatives and assessed their biological activity against FP2. Accordingly, the S2 pocket is accessible for different substituents of 5- or 6-membered aromatic systems such as pyridin-3-yl, 4-methoxyphenyl, 2-methoxyphenyl, 3,4-dimethoxyphenyl, furan-2-yl, and thiophen-2-yl. Also, loss of activity was observed when the 2-aminosobutyric acid substituted the 1-aminocyclohexanecarboxylic acid as a center core. Most of the derivatives exhibited high selectivity against FPs versus cathepsin B and L, as well as against the serine protease α-chymotrypsin, making them interesting targets for further optimization studies toward the discovery of new potent antimalarials. Compound 25, one of the most active inhibitors of the series against FP2 action is displayed in (Fig. 20).

Fig. (20). Compound 25, one of the best inhibitors designed by Diederich and co-workers [103] with a Ki value against FP2 of 1.2 μM.

Most recent examples of peptidyl-based structures as potential FP inhibitors include those reported by Gomes and co-workers [105]. They designed and synthesized a series of cinnamic acid/4-aminooquinoline derivatives as potential inhibitors of falcipain action and found the unsubstituted cinnamic acid derivative 26 (Fig. 21) to inhibit not only FP2 action but also the parasite growth in vitro in the micromolar range. The retro enantio dipeptide used to link the 4-aminooquinoline and the respective cinnamic acid was demonstrated to provide antimalarial activity. Interestingly, using the natural counterparts of the amino acids in the linker
dipeptide was detrimental to compound inhibitory activity against FP2 and also resulted in a decrease in the antiplasmoidal activity. Furthermore, some of the compounds from the series exhibited activity against hemozoin biocrystallization and FP2 making them good leads toward the development of potential dual action antimalarials.

Fig. (21). The most active FP2 inhibitor among the new series proposed by Gomes and co-workers [105].

All the above examples demonstrate that, with the exception of vinyl sulfones unveiled by Rosenthal’s group, compounds able to irreversibly inhibit FP and display nanomolar activities against Pf have been hard to find. Hence, several groups have developed novel structures where the vinyl sulfone moiety is conserved in an attempt to ensure FP irreversible inhibition. An example of such strategy is that of Zappalà’s team [92, 93]. These authors synthesized a series of Cys protease inhibitors based on the 1,4-benzodiazepine scaffold, which is known to act as a good mimetic of β-turns [113], the structural motif postulated as the biological active form of the D-Ser-Gly peptide [114]. Furthermore, benzodiazepines are also known to enhance oral bioavailability and also to increase stability toward premature proteolytic degradation by enzymes. In this connection, Zappalà and co-workers [93] focused their attention on the synthesis of two different series both including a vinyl sulfone in the P1’ site (Fig. 22). One series contained hPhe in P1, a residue known to increase the inhibitors potency against FPs, and the second series included Gly in P1 to assess the relevance of the amino acid side chain for enzyme recognition. All of the vinyl sulfone inhibitors displayed activity against the parasite higher than 9.1 μM. As expected, compounds with hPhe residue in P1 site were generally better inhibitors of parasite development than compounds containing a Gly. Compounds 27a and 27b (Fig. 22) presented good FP2 inhibitory activities and also the highest antiplasmoidal activities of the corresponding series [93]. In addition, the compounds were tested against cathepsin B and L and found to be weak inhibitors of these two enzymes, which can be considered a promising result regarding their selectivity toward parasite versus host Cys proteases. Thus, these compounds can be considered as leads for the development of new antimalarials.

Following, Zappalà’s group reported a series of compounds synthesized based on the same 1,4-benzodiazepine scaffold with the aim of improving the pharmacological properties of this type of irreversible inhibitors [92]. In that work, they described a highly potent and selective derivative with a vinyl ester warhead 28 (Fig. 23), which exhibited the highest potency with an activity two times higher than that of the standard E-64 ($k_{\text{eff}}=1586000 \text{ M}^{-1} \text{ min}^{-1}$) and the highest enzymatic affinity ($K_i=17 \text{ nM}$) among the series. Although 28 displayed a $K_i=17 \text{ nM}$, it did not result as active in vivo, supposedly, due to the difficulty for the derivative to cross the biological membranes of the parasites and reach FP2.

Fig. (22). FP2 inhibitors, designed by Zappalà’s team, combining a benzodiazepine inhibitors and a vinyl sulfone warhead [93].

Fig. (23). Most potent compound of the series 28 reported by Zappalà and co-workers [92].

The unmatched relevance of the dipeptidyl vinyl sulfone warhead for FP inhibition has been also recognized in research works where this moiety has been combined with other antimalarial scaffolds, e.g., artemisinin, to produce multi-target antimalarial molecules 29 (Fig. 24) [115].

Fig. (24). Artemisinin-dipeptidyl vinyl sulfone hybrids 29 developed by Moreira’s team [115].

2.2.1.2. Reversible Inhibitors

An example of reversible inhibitors was reported by Zappalà’s team [104]. These authors synthesized a series of
Cys protease inhibitors based on the 1,4-benzodiazepine scaffold linked to a C-terminal aspartyl aldehyde building block that would inhibit the enzyme by forming a covalent reversible bond. All of the designed benzodiazepine derivatives presented activity against FP2 with IC_{50} values between 8 and 26 μM. The derivative of compound 30 obtained by introducing a phenyl group at R position (Fig. 25) was the least potent inhibitor of the series with an IC\textsubscript{50} of 21.54 μM. The introduction of an electro-withdrawing or electron-donating group to the para position of the phenyl ring in the general structure 30 (Fig. 25) increased the activity against the enzyme. An enhancement of the activity against FP2 was also observed by the insertion of a methyl or a trifluoromethyl group to the p-chlorophenyl ring (c.f. compound 31 in Fig. 25). To evaluate the selectivity of the compounds, the derivatives were tested against a panel of active recombinant human caspases (i.e., caspases 1-9) and the molecules displayed inhibitory activity up to 50 μM.

![Fig. (25). General structure 30 for the design of benzodiazepine inhibitors with a P1-aspartyl aldehyde moiety and compound 31, one of the best inhibitors of the series [104].](image)

Other examples of reversible inhibitors include the ones reported by O’Neill and co-workers [90]. They studied examples of FP2 and FP3 inhibitors that do not have an αβ-unsaturated system included in the structure and inhibit FPs in a reversible manner. They were able to synthesize two molecules, namely, compounds 32 and 33 in (Fig. 26), bearing an aldehyde moiety which displayed activity against FP2 and FP3 in the low nanomolar range, and also expressed activity versus the 3D7 strain of Pf. Both molecules contained residues Leu and homoPhe, the key amino acids previously identified by Rosenthal and co-workers [90], and were found to possess a good enzyme fit according to docking studies.

Other peptidyl reversible inhibitors of falcipains include peptidyl aldehyde and α-ketoamide derivatives developed by Rosenthal and co-workers [91]. They evaluated FP2 inhibition and antimalarial activity of the series and found most of the compounds to inhibit FP2 activity in nanomolar range. Due to the high sequence similarities between FP2 and FP3, they decided to test the compounds also against FP3 and activity against both falcipains resulted to be similar, suggesting that a single small specific protease inhibitor could be enough to inhibit both falcipains and consequently, inhibit hemoglobin degradation leading to parasite death. As seen previously in other works, the compounds which displayed higher activity against the protease had a Leu in P2 site contrary to those who had a Phe in P2. Inhibition of FP2 was always accompanied by swollen and darkly stained parasitic FV, an experimental finding that agrees with blocking of hemoglobin degradation [116]. Since new ideal antimalarials should be active against all Pf strains, Rosenthal and co-workers decided to test the three best inhibitors of the series against other five Pf strains. They found the three compounds to inhibit more or less equally each strain and that there was no evidence that the activity against multidrug-resistant strains differed from the activity against broadly sensitive strains. The morpholino-carbonyl-leucine-homophe-nylalanine aldehyde 34 (Fig. 27) exhibited excellent activity against FP2 and parasite development. Rosenthal’s team assessed the activity of the P. vinckei-mice and the compound 34 showed antimalarial activity when it was administered intraperitonially but this activity was fairly low. They inferred that compound 34 activity could be mainly due to the short life of the compound. Thus, due to the poor pharmacokinetik of the compound to provide consistent levels in blood and to improve drug delivery, mice were dosed by a subcutaneous infusion pump continuously. However, the antimalarial activity continued to be modest and the mice eventually died by the infection.

![Fig. (26). FP2/3 reversible inhibitors developed by O’Neill and co-workers [90]. Mu=Morpholine urea; Cbz=Benzylxocarbonyl.](image)

![Fig. (27). Most active peptidyl reversible inhibitor 34 developed by Rosenthal and co-workers [91].](image)
2.2.2. Non-Peptidic Inhibitors

In an effort to develop non-peptidic FP2 inhibitors, Zap-palà’s group reported a series of 1-aryl-6,7-disubstituted-2H-
isoquinolin-3-ones which were synthesized and tested
against FP2, as well as against cultured Pf strain FCBR para-
sites. Most compounds presented activity against the enzyme
though there were not selective. The most potent com-


gound of the series 35 (Fig. 28) displayed a $K_i=2.3 \mu M$ against FP2
emphasizing the importance of hydrophobic and bulky
groups at position 6 and 7 of the isoquinoline scaf-
dfold [117]. Further work is required in order to improve the sel ectivity of
these compounds because they resulted to be slightly more
active against human cathepsins than FP2.

![Fig. (28). Most potent compound 35 of the series reported by Zap-palà and co-workers [117].](image)

Several other non-peptidic small inhibitors of FPs have
been found [118-121], some of them through computer-
based drug design and virtual screening studies, for instance,
compounds 36-39 (Fig. 29) [94-96, 120]. Recently, Avery
and co-workers [120] identified a total of 28 non-peptidic
low micromolar inhibitors of FP2 and FP3 and elaborat-
ed SAR for each series. Some of the complex trends in the SAR
could be explained by the energetical changes associated
with the displacement of water molecules upon ligand bind-
ing, and other could be related with poor chemical reactivity
of the reactive centers of these compounds. The motivation
to use water energetical changes was due to the fact that the
SAR could not be explained by none of the following de-
scriptors: hydrogen bonds, van der Waals, and electrostatics
interactions, or by ligand strain, or even by using traditional
computational approaches such as molecular mechani-
cal/energetics/ generalized Born surface area (MM-GBSA), docking
scores, and molecular mechanics interaction energies. Some
of the molecules identified in this study also inhibited para-
site growth in culture.

![Fig. (29). Non-peptidic FP inhibitors developed by Avery and co-
workers [120].](image)

Some hybrid drugs, capable of targeting two processes in
the malaria life cycle where one of the targets is FP2 inhibi-
tion, can fall into the category of non-peptidic small inhibi-
tors [122, 123]. For instance, in 2010, Chibale’s group iden-
tified chalcone-chloroquine hybrid 40 as a promising antimal-
arial (Fig. 30) [122, 123]. Compound 40 and related hybrid
structures were synthesized and showed to be active against
Pf. The molecules were tested as inhibitors of both FP and
hemozoin formation, and some of them were found to be
active in the micromolar range against FP. However, com-

![Fig. (30). Hybrid drug 40 designed to include FP inhibition (α,β-
unsaturated system as Michael acceptor) in its mode of antimalarial
action [122, 123].](image)

some hybrid drugs, capable of targeting two processes in the malaria life cycle where one of the targets is FP2 inhibition, can fall into the category of non-peptidic small inhibitors [122, 123]. For instance, in 2010, Chibale’s group identified chalcone-chloroquine hybrid 40 as a promising antimalarial (Fig. 30) [122, 123]. Compound 40 and related hybrid structures were synthesized and showed to be active against Pf. The molecules were tested as inhibitors of both FP and hemozoin formation, and some of them were found to be active in the micromolar range against FP. However, compound 40 and related structures did not show correlation between FP inhibition and parasite growth in culture. Still, there was consistent correlation between in vitro antimalarial potency and inhibition of hemozoin formation, suggesting that this could be the primary mechanism of their antimalarial activity.

![Fig. (30). Hybrid drug 40 designed to include FP inhibition (α,β-
unsaturated system as Michael acceptor) in its mode of antimalarial
action [122, 123].](image)

Other non-peptidic inhibitors include structural analogs of triazole (Fig. 31) and benzothiazole (Fig. 32) cores showing moderate inhibition of falcipains, reported by Avery and co-workers [124]. In the triazole series, in an effort to optimize the activity of compound 41 against FP2, only few tria-
zoles with general structure 42 were found to display activity against FP2 but none was more active than the parent com-
compound. The low activity of these derivatives might be ascribed to the unsuitable or shorter hydrophobic R groups used for this family as compared to the parent compound 41 that had an IC\textsubscript{50} value of 2.2 \mu M. Indeed, molecules with bulkier R groups were the ones exhibiting modest activity against FP2.

![Fig. (31). General structure used to design triazoles by Avery’s team [124].](image)

In the benzothiazole series, compounds 43 and 44 were predicted by docking studies to interact with the polar residues of the S2 pockets of FP2 and FP3; these compounds inhibited both FP2 (IC\textsubscript{50} between 12.22-12.75 \mu M) and FP3 (IC\textsubscript{50} between 13.77 and 14.94 \mu M), but also displayed activity against homologous mammalian cysteine proteases lacking polar residues, suggesting their low selectivity. Only compounds 44 and 45 inhibited the growth of W2 strain of Pf, with IC\textsubscript{50} values of 2.08 \mu M and 4.65 \mu M, respectively.

![Fig. (32). Benzothiazole based inhibitors as Plm II designed by Avery and co-workers [124].](image)

Other non-peptidic FP inhibitors include heteroaryl nitrites. For instance, Fiandor and co-workers [118] explored the pyrimidinenitrile scaffold for FP inhibition: optimization studies led to the discovery of 2-cyano-5-chloropyrimidines and 2-cyano-5-bromopyrimidines as promising inhibitors of the enzymes. Although some compounds inhibited falcipains in the subnanomolar and low nanomolar ranges, they only inhibited parasite growth in the micromolar range. However, introduction of a protonable amine resulting in analogs such as compound 46 (Fig. 33) led to an increase in antiplasmodial activity.

![Fig. (33). One of the best 2-cyano-5-bromopyrimidines, 46, developed by Fiandor’s team: FP2 IC\textsubscript{50}=0.5 nM, FP3 IC\textsubscript{50}=3 nM and Pf W2 IC\textsubscript{50}=1 nM [118].](image)

Organometallic reversible FP inhibitors have also been proposed, namely, a series of gold derivatives reported by Messori and co-workers [125]. They found gold complexes capable of pronounced and reversible inhibition of FP2 in the micromolar range. Still, they were unable to establish a correlation between enzyme inhibition and antiplasmodial activity. Compound 47 (Fig. 34) displayed the lowest \(K_{i}\) (1.4 \mu M) of the series with an IC\textsubscript{50} of 5.11 \mu M against parasite growth.

![Fig. (34). Gold derivative 47, with micromolar activity against parasite development [125].](image)

More recently, Gomes and co-workers [105] developed non-peptidic inhibitors of FP2 including a cinnamoyl moiety. They synthesized a series of cinnamic acid/4-amino quinolines conjugates 48 (Fig. 35) with FP2 inhibitory activity in the micromolar range. By analyzing the binding modes of 48 docked to FP2, those authors observed that most of the compounds present the cinnamoyl group at the S2 cavity and having the vinyl group relatively close to the active residue of the cysteine protease. However, none of these inhibitors displayed antiplasmodial activity, probably due to low permeation into the infected red blood cell or into the parasitophorous vacuole inside it.

The combination of the inhibitory activity results regarding all falcipain inhibitors reported so far with the recent availability of crystal structures of both FP2 and FP3, will
strongly help and support all the medicinal chemists currently involved in the discovery of potent falcipain inhibitors as promising molecules to combat malaria disease.

Fig. (35). Best inhibitor of the cinnamic acid derivatives reported by Gomes’ team [105].

2.3. Falcilysin

The zinc metalloprotease falcilysin (FLN) is another protease that participates in the hemoglobin degradation process, specifically, in the hydrolysis of short globin peptides produced by FP-mediated proteolysis of globin [75, 126]. FLN is highly active at acidic pH, which is consistent with its role in the FV, but it is also active at neutral pH, although with a different substrate specificity. At neutral and acidic pH this protease prefers to cleave sites in which P1’ residues are bulky hydrophobic amino acids and has strong preference for methionine at P3’; however, in what concerns P2’ residues, FLN favors hydrophobic residues at pH 5.2, whereas at pH 7.2 it has stronger preference for Arg. Finally, at P4’ and P5’, FLN prefers acidic residues at acidic pH but is less selective at neutral pH. Studies by Goldberg and co-workers [127] revealed that FLN is not only present in the FV but also in association with vesicular structures elsewhere in the parasitophorous vacuole, suggesting an expanded role for this protease in Pf biology. Despite some chelators, such as 1,10-phenanthroline [128] presented some activity toward this protease, no specific inhibitor was found so far [127-129].

2.4. Dipeptidyl Amino Peptidase 1

Once FLN converts short globin polypeptides into oligopeptides consisting of 5–10 amino acids, dipeptidyl amino peptidase 1 (DPAP1), a lysosomal exopeptidase, sequentially cleaves oligopeptide substrates from their N terminus into dipeptides [87, 130]. Attempts to disrupt the DPAP1 gene have been unsuccessful, which suggests that the enzyme makes an important contribution to hemoglobin catabolism during the intraerythrocytic cycle [131]. While DPAP1 is one of the three related DPAP enzymes encoded in the parasite genome, current evidence suggests that only DPAP1 resides in the food vacuole [130]. DPAP1 is also potentially a good drug target because the closest human homologue is cathepsin C, which is not essential in mammals [130, 132, 133]. DPAP1 differs from other endopeptidases in that they recognize the N-terminal free amine of substrate proteins and cleave N-terminal dipeptides [134]. Structurally, its active site is relatively small compared to that of other endopeptidases and its substrate specificity is mainly dictated by the N-terminal residue P2 [134]. This smaller active site is more amenable to binding of bioactive molecules with low molecular weight and should be more suitable for computational docking and in-silico design of inhibitors.

Although DPAP1 is a fairly new target, some inhibitors of this protease have been already developed: Bogyo and co-workers [134] screened several non-peptidic scaffolds from a large library of small compounds that was initially designed to target cysteine cathepsins in mammals, and found stable covalent DPAP1 inhibitors 49 (Fig. 36) which kill Pf at low nanomolar concentrations.

Bogyo findings demonstrated that inhibition of DPAP1 by small compounds results in an immature trophozoite and, consequently, parasite death. This strongly suggests that development of DPAP1 inhibitors may be a key strategy toward malaria chemotherapy.

Fig. (36). Best DPAP1 inhibitor found by Bogyo’s team [134].

2.5. Exoaminopeptidases

Pf contains nine exoaminopeptidases [135], four of which are methionine aminopeptidases (MetAPs). MetAPs catalyze the removal of N-terminal methionine during protein synthesis [129, 136] and their inhibition can impair Pf growth in culture [24, 129, 137, 138]. The remaining five Pf exoaminopeptidases include an aspartic aminopeptidase, a prollyl aminopeptidase, a post-prolyl aminopeptidase, a leucine aminopeptidase, and an alanine aminopeptidase. These five aminopeptidases are considered to be potential targets for inhibition of hemoglobin degradation. Neutral alanine and leucine metallo-aminopeptidases [139], involved in the generation of free amino acids from the previously generated dipeptides, are the pivotal Pf exoaminopeptidases. Bestatin 50 (Fig. 37) has been found to inhibit both the activity of these enzymes and malaria parasite growth in vitro [36].

Fig. (37). Bestatin 50, a reported inhibitor of Pf Leu and Ala aminopeptidases [36].

2.5.1. M1 Alanine Aminopeptidase

P/A-M1 (EC number: 3.4.11.2) is a plasmodial zinc aminopeptidase that belongs to the M1-family which displays its activity at pH 7.4 and has a broad substrate spectrum [140, 141]. P/A-M1 is found in both trophozoites and
schizonts, being involved in hemoglobin degradation and erythrocyte reinvasion [142]. Therefore, design and synthesis of potential inhibitors of the PfA-M1 activity can lead to the discovery of new antimalarials [141, 143, 144]. For example, Deprez-Poulain’s group first identified three non-peptidic inhibitors of this protease, and further optimized their results, and found the desired specificity against mammalian neutral aminopeptidases (APN), the host M1 family prototype [141]. The compounds were based on malonic hydroxamic template 51 (Fig. 38) and the influence of the malonic substituents on the activity against PfA-M1 was analyzed: compounds bearing a benzyl or a m-phenoxymethyl benzyl group in R1 presented higher activity compared to those with an isobutyl substituent in that position. Binding of hydroxamate to the Zn\(^{2+}\) is expected to be the main anchoring point of the potential inhibitor. Therefore, substituents on R1 position should discriminate those molecules which will bind to the enzyme from those which will not. Compounds bearing a benzyl amine in R2 position, when presenting a fluorine atom in the para position or a methyl group on the ortho position, expressed strong increase on activity. To remove the chiral carbon, the same authors synthesized unsaturated analogues of 51, and found that a Z-configuration 52 led to increased activity, whereas an E-configuration had the opposite effect. Additionally, they found that cyclization of 52 to produce 53 was detrimental for enzyme inhibitory activity.

![Fig. (38). PfA-M1 inhibitors developed by Deprez-Poulain’s group](attachment:image)

In addition, Deprez-Poulain’s group has also developed quinoline inhibitors of PfA-M1 [145]. They discovered that the presence of a Zn-chelating group is essential for the activity. They also found that the hydroxamate group is more efficient than a carboxylic acid group. The three PfA-M1 inhibitors are also inhibitors of the parasite growth and the quinoline moiety allows these compounds to also inhibit heme biocrystallization. The best inhibitor 54 (Fig. 39) of the synthesized compounds exhibited IC\(_{50}\) values of 0.854 \(\mu\)M and 0.317 \(\mu\)M against PfA-M1 or parasite growth, respectively.

Recently the X-ray structures of PfA-M1 in the apo form, in complex with bestatin (50, Fig. 37), and with a phosphine-based inhibitor, hPheP(CH\(_2\))Phe (55, Fig. 40), originally designed for the cytosol leucine aminopeptidase [146], were determined by Dalton and co-workers (PDB codes: 3EBG, 3EBH, 3EBI) [143]. The availability of these crystal structures will trigger the explanation of different affinities presented by different inhibitors allowing the early phase of inhibitors optimization.

![Fig. (39). The best quinoline-based inhibitor of PfA-M1 developed by Deprez-Poulain’s group](attachment:image)

### 2.5.2. M17 Leucine-Aminopeptidase

There is an overexpression of M17 leucine aminopeptidases (EC number: EC 3.4.11.1) in the malaria parasite; actually, the expression level of M17-family is ~18-fold higher than the M1-family [147, 148]. Leucine aminopeptidase (LAP) is an exopeptidase that has a preference for substrates which contain leucine or alanine in the N-terminus. Plasmodial LAP (P/LAP) exhibits its optimal activity at pH 7.2 and is inactive below pH 6, which is consistent with no activity in the FV. Apart from bestatin 50, other inhibitors of M17 leucine aminopeptidases have been reported [149, 150]. For instance, Gardiner and co-workers reported in 2007 a series of phosphinate dipeptide analogues as potential P/LAP inhibitors [150]. By screening different phosphinate analogues of dipeptides against functionally recombinant P/LAP, those authors were able to identify four compounds as potent inhibitors of the parasite P/LAP activity and from those four molecules, compounds 55, 56 displayed in (Fig. 40) presented the highest inhibitory effect against P/LAP. These two compounds resulted to be even more active than bestatin 50 against P/LAP and also showed activity against malaria parasite growth in culture in the low micromolar range.

Docking of compound 55 against the enzyme showed that an amino group and two oxygen atoms are able to interact with the two zinc metal ions in the active site of P/LAP. In addition, the hydrophobic S1 in P/LAP can accommodate the hPhe of ligand 55. The affinity of compound 56 to P/LAP was significantly similar to that of 55 because the additional hydroxyl group from the Tyr residue is solvent exposed and does not promote additional interactions with the enzyme. Molecular docking studies showed that the phosphinic moieties in both 55 and 56 are able to form unique contacts with the metals ions and residue Lys386 of the enzyme’s active site. These interactions do not occur with the corresponding fragment in bestatin and, therefore, can be accounted for the high binding affinity of 55 and 56 over bestatin 50 for
P/LAP. Compound 55 was taken further to in vivo analysis using a non-lethal murine malaria model and the inhibitor 55 reduced the parasitemia of mice infected with *P. chabaudi* (reduction of parasite burden: 92%) [150].

![Fig. (40).](image)

3. CONCLUDING REMARKS

Since the unveiling of the *Pf* genome, the number of targets to inhibit malaria parasite activity within host erythrocytes has increased, and there has been a great progress toward the development of new antimalarials. Still, malaria continues to be one of the most common infectious diseases and a burden on developing nations, mainly due to the fast growing resistance of parasites to existing drugs. Plasmodial proteases which participate in the hemoglobin degradation have been found to be potential antimalarial targets since inhibition of this proteolytic process will impair parasite growth by starvation, and also would increase colloid-osmotic pressure within the infected erythrocyte causing its premature lysis. This review provides a general overview of the main proteases which participate in hemoglobin degradation currently being studied to, in time, suppress morbidity and mortality caused by malaria and prevent the progression of uncomplicated malaria to potentially fatal disease.

CONFLICT OF INTEREST

The author(s) confirm that this article content has no conflicts of interest.

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