A Class of Valuable (Pro-)Activity-Based Protein Profiling Probes: Application to the Redox-Active Antiplasmodial Agent, Plasmodione

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ABSTRACT: Plasmodione (PD) is a potent antimalarial redox-active drug acting at low nM range concentrations on different malaria parasite stages. In this study, in order to determine the precise PD protein interactome in parasites, we developed a class of (pro-)activity-based protein profiling probes (ABPP) as precursors of photoreactive benzophenone-like probes based on the skeleton of PD metabolites (PDO) generated in a cascade of redox reactions. Under UV-photoirradiation, we clearly demonstrate that benzoylmenadione 7 produces the 3-benzoylmenadione probe 9, allowing investigation of the proof-of-concept of the ABPP strategy with 3-benzoylmenadiones 7–10. The synthesized 3-benzoylmenadiones, probe 7 with an alkyne group or probe 9 with -NO2 in para position of the benzoyl chain, were found to be the most efficient photoreactive and clickable probes. In the presence of various H-donor partners, the UV-irradiation of the photoreactive ABPP probes generates different adducts, the expected "benzophenone-like" adducts (pathway 1) in addition to "benzoxanthone" adducts (via two other pathways, 2 and 3). Using both human and Plasmodium falciparum glutathione reductases, three protein ligand binding sites were identified following photolabeling with probes 7 or 9. The photoreduction of 3-benzoylmenadiones (PDO and probe 9) promoting the formation of both the corresponding benzoxanthone and the derived enone could be replaced by the glutathione reductase-catalyzed reduction step. In particular, the electrophilic character of the benzoxanthone was evidenced by its ability to alkylate heme, as a relevant event supporting the antimalarial mode of action of PD. This work provides a proof-of-principle that (pro-)ABPP probes can generate benzophenone-like metabolites enabling optimized activity-based protein profiling conditions that will be instrumental to analyze the interactome of early lead antimalarial 3-benzoylmenadiones displaying an original and innovative mode of action.

KEYWORDS: activity-based protein profiling, antimalarial, 3-benz(o)ylmenadione, CuAAC, electrophile, photoaffinity labeling, photoredox, quinone

INTRODUCTION

To decipher drug modes of action (MoA), chemical strategies for functional proteomics have been developed in the recent years with the activity-based protein profiling (ABPP) being one of the most specific. This unbiased and alternative methodology to identify drug or drug metabolite interactors in diverse organisms has successfully detected protein partners of miscellaneous biomolecules (see the pioneering work from Cravatt et al. and then from Bogyo et al., with the first reports about serine hydrolase inhibitors). ABPP allows monitoring and dissecting a drug interactome from complex proteomes in their native forms. This is achieved by the design and synthesis of small drug-activity-based probes that can react with the protein targets. The selective separation of the drug-protein adducts from the whole proteome is made possible by the Cu(I)-catalyzed alkyne–azide cycloaddition (CuAAC) “click” reaction, also known as the seminal Huisgen reaction adapted for biological chemistry and proteomics purposes. The ABPP probe is based on three essential functional chemical elements: (1) a recognition group—drug/metabolite that has affinity to specific proteins defined as interactome; (2) a reactive group—either electrophilic or photoreactive feature that favors cross-linking or covalent binding of the probe to the protein targets. The selective separation of the drug–protein adducts from the whole proteome is made possible by the
target(s); (3) a reporter group/tag (e.g., alkyne or azide) enabled to react in the click reaction with a partner (e.g., azide or alkyne) that is functionalized either by a fluorophore for visualization of the drug−protein adducts or a affinity chromatography tag for enrichment and identification of the adduct. An additional advantage in the ABPP field is the enlargement of the variety of chemical probes that trap representatives of various enzyme classes and can be utilized in
proteome studies. In the case of the flavin-dependent oxidoreductase family, the design of clickable ABPP probes was limited to the cytochrome P450, 2-oxoglutarate oxidases, and amine oxidases. According to the literature, several proteomics studies have been conducted with ABPP probes for the detection of drug targets in P. falciparum.

NAD(P)H-dependent flavoenzymes from the malarial parasites have been proposed as possible targets of plasmoidine, an early antimalarial lead compound (PD, 3-[(trifluoromethyl)benzyl]-menadione, I, Figure 1). Initial studies have focused on the chemical reactivity of the 3-benzoylmenadione core, and of its key putative metabolites, the 3-benzoylemenadiones, which were shown in vitro to act as effective subversive substrates of recombinant glutathione reductases from human (hGR) and P. falciparum ( PfGR). GR is a homodimeric NADPH-dependent FAD-containing enzyme ( GR; EC 1.8.1.7) that belongs to the family of NADPH-dependent oxidoreductases. GR catalyzes the reduction of glutathione disulfide (GSSG): NADPH + H+ + GSSG ⇌ NADP+ + 2 GSH (eq 1). According to previous studies, suicide-substrates such as fluoromethyl menadione derivative, inactivates GSSG reduction by hGR but not naphthoquinone reduction, thus suggesting that naphthoquinone reduction occurs at a different site than GSSG reduction, possibly near the flavin, close to the NADPH binding site, as previously postulated. Subversive substrates inhibit GSSG reduction activity because they are reduced by the NADPH-reduced flavin enzyme species, thus preventing electrons to flow normally from NADPH to GSSG. In the presence of natural oxidizers, for example, oxygen or methemoglobin(Fe3+) (metHb), naphthoquinone reduction was demonstrated to be reversible for PD and its key metabolite, the 3-benzoylemenadione (PDOred, compound 2), starting a redox-cycling process (Figure 1A). Reduced benzoylemenadiones can efficiently transfer one electron to metHb, and the redox cycle constantly regenerates the benzoylemenadione under its oxidized form at the expense of the NADPH pool. MetHb is a critical nutrient for Plasmodium crucial for its growth (e.g., at the trophozoite stage) while Hb(Fe2+) is not digestible. The shift in metHb/Hb(Fe2+) balance during redox-cycling results in metHb depletion and parasite growth arrest. From PD, the NADPH-dependent oxidoreductase-promoted redox-cycling also produces a continuous flux of reactive oxygen species (ROS) and toxic metabolites, including the reduced 3-benzoylemenadione (PDOred, compound 3) (i.e., via 1-electron transfer) along with hemoglobin degradation catabolites identified as membrane-enriched hemichromes. The latter are known to act as biomarkers of red blood cell (RBC) senescence and to trigger early phagocytosis by macrophages. Hence, PD activation via PDO-mediated redox-cycling most likely results in the specific removal and clearance of the parasitized RBCs (pRBC).

Furthermore, during metHb digestion, toxic heme is released into the acidic food vacuole of the parasite. To detoxify free heme, the parasite converts it into a nontoxic insoluble hemozoin biocystal. We previously proposed that PD bioactivation in pRBCs, possibly by GR, generates a key metabolite—the benzoxanthone (PDO-BX) (Figure 1A) via a cascade of redox reactions and oxidative phenolic coupling. In turn, PDO-BX can firmly interact with free heme and is thus suggested to prevent heme crystallization leading to parasite death.

Finally, in yeast, the mitochondrial type II NADPH-dehydrogenase Nde1, was found to be the main target responsible for PD redox-cycling, with GR and two other oxidoreductases (Mcr1 and Lpd1) being minor targets. Taken together, these observations and the current model for PD MoA suggest that, once generated, PD metabolites could (i) redox cycle with several oxidoreductases, which may vary according to parasite developmental stage, generating oxidative stress; and/or (ii) disturb key parasite processes such as hemozoin formation. Additionally, the abundance of proteins expressed in parasites is variable and depends on the parasite stages. Thus, during any ABPP study, actual drug targets expressed in traces would be difficult to distinguish from unspecific labeled but abundantly expressed proteins recovered in the HPLC MS/MS analysis. The focus of the present study was therefore to design a series of relevant and specific PD-ABPP probes, to define standardized conditions for their use and establish a proof-of-concept of their application with isolated proteins such as hGR and PfGR as models (Figure 1C). Here, for the first time, we report that 3-benzoylemenadiones are photoreactive and, as (pro-)activity-based protein profiling probes ((pro-)ABPP), can be used for ABPP applications.

The 3-benzoylemenadione probe generates a benzophenone-like moiety upon photoreduction, a step that mimics the redutive bioactivation drug pathway catalyzed by a flavoenzyme in the living cell. Diversely substituted benzophenone-like and BX adducts were produced in the presence of different partners via original photoredox pathways that have not been previously described. The successful photoaffinity labeling of both GRs not only allowed the identification of naphthoquinone binding sites in GR structure but also revealed an alkylation process of the toxic heme by PDO-BX, generated upon PD redox-cycling with hGR, which is likely a relevant event contributing to PD MoA.

**RESULTS AND DISCUSSION**

**Design of 3-Benzoylmenadiones as Photoreactive Probes**

Our original strategy for designing the PD-ABPP is leveraged from the postulated MoA of PD. PD was suggested to act as a prodrug generating in situ a key metabolite, PDO, upon PD bioactivation (i.e., benzyl oxidation) (Figure 1A). Interestingly, PDO possesses in its structure the 2-benzoyl-1,4-naphthalenedione group that could behave as a 2-benzonaphthophene precursor. Therefore, we assumed that the redox-active PD-derived benzoylemenadione, under its reduced state, might have an intrinsic photoreactive benzophenone-like structure per se and thus permit the spontaneous covalent trapping of targets upon photoirradiation. Such probes might show high spatiotemporal control of targeted enzyme recognition/alkylation and drastically lower nonspecific binding. Interestingly, no additional bulky photoreactive group was introduced into the PD metabolite structure. Furthermore, the newly designed ABPP probes 7–11 (Figure 1B) were functionalized in the benz(o)yl chain, by a reporter alknye group known to bring minimal structural and electronic perturbation. To validate the hypothesis of the intrinsic photoreactivity properties attributed to the benzophenone-like structure, we first studied the 3-benzoylemenadione derivatives in model photochemical reactions. We compared the photoreactivity of the previously reported PD analogue 3-benzoylemenadione...
5−6 pair17 (Figure 1B) upon photoirradiation at 350 nm (Figure 2) in comparison with benzophenone (Figure S1) to evaluate whether the keto group of the benzoyl chain is essential for photoreaction. For this, we used N-acetyl-methionine methyl ester (N-Ac-Met-OMe, shortened as nMet) as a partner model, in accordance with previous studies demonstrating that methionine and its N-Ac-Met-OMe derivative are predominantly alkylated at the side-chain in α-position to the sulfur.24,25 The products of the photoreaction were analyzed by field desorption-mass spectrometry (FD-MS). As observed in Figure 2A, the insertion product of the 3-benzoylmenadione derivative 6 and nMet displayed a mass

Figure 2. Mass spectrometric analysis of the photochemical reaction mixtures. Field-desorption mass spectrometry (FD-MS) analyses of the photochemical reaction mixtures of (panel A) the 3-benzoylmenadione 6 or (panel B) the 3-benzylmenadione 5 derivatives, in the presence of the diprotected methionine nMet.

Scheme 1. Synthesis of the 3-Benzoylmenadiones 7−10 (Paths A and B) through the Friedel−Crafts Reaction Variant31 and the 3-Benzylmenadione 11 (Path C) through the Kochi−Anderson Reaction32,24

“Reaction conditions: (i) 1. SnCl2 cc HCl, EtOH, rt, 2 h, 2. Me3SO4 acetone, KOH, MeOH, 60 °C, 4 h; (ii) TiOH/TFAA, DCM, and A. 4-iodobenzoic acid 7a, or 3-iodo-4-(trifluoromethyl)benzoic acid 8a. B. 4-nitro-3-fluorobenzoic acid 9a, or 3-fluoro-4-(trifluoromethyl)benzoic acid 10a, 0 °C then rt, 16 h; (iii) Cul, Pd[PPh3]2Cl2, NEt3, rt, 16 h, HC ≡ C-TMS; (iv) TBAF, THF, rt, 1.5 h; (v) CAN, H2O/MeCN, rt, 3 h; (vi) K2CO3 DMF, propargylic alcohol, 60 °C, 24 h; (vii) 4-iodophenylacetic acid, AgNO3 Na2S2O8 MeCN/H2O, reflux, 4 h.

5−6 pair17 (Figure 1B) upon photoirradiation at 350 nm (Figure 2) in comparison with benzophenone (Figure S1) to evaluate whether the keto group of the benzoyl chain is essential for photoreaction. For this, we used N-acetyl-methionine methyl ester (N-Ac-Met-OMe, shortened as nMet) as a partner model, in accordance with previous studies demonstrating that methionine and its N-Ac-Met-OMe derivative are predominantly alkylated at the side-chain in α-position to the sulfur.24,25 The products of the photoreaction were analyzed by field desorption-mass spectrometry (FD-MS). As observed in Figure 2A, the insertion product of the 3-benzoylmenadione derivative 6 and nMet displayed a mass
peak at \(m/z = 526.2\). Under the same photoirradiation conditions, the photoreactive benzophenone generated the insertion product with nMet as attested by the presence of mass peaks at \(m/z = 387.2\) (M\(^+\)) and 369.2 (M\(^+\)_H\(_2\)O) (Figure S1). No major insertion product was observed for the 3-benzoylmenadione derivative 5 (Figure 2B), demonstrating that the 3-benzoyl chain is essential for the photoreactivity of the benzoylmenadione derivative 6.

Interestingly, probe 6 appears to be photochemically reactive per se, even in the absence of a prereduction step in the presence of the NADPH/GR. The 3-benzoyl-dihyronaphthoquinone is likely generated by photoreduction upon photoirradiation, thus allowing formation of the insertion product. At first glance, we neglected the small peaks appearing at \(m/z = 321.1\) and 323.1 suggesting the formation of trace photoproducts upon photoirradiation of the 3-benzylmenadione 5, which were finally attributed to both oxidized and reduced species of the 3-benzylmenadione 6 (Figure 2, black box). This observation was confirmed later when we used the most photoreactive 3-benzylmenadione ABPP probe 11 per se to investigate the generation of photoproducts upon UV-photoirradiation.

Using FD-MS under the same experimental conditions with 3-benzoylmenadiones, we were able to observe the insertion product of the benzoylmenadione 6 with a p-nitrobenzoyl derivative but not with benzylmenadione 12 with a carboxylic acid group in para- to the benzoyl ring (data not shown). This result might be explained by the fact that the carboxylate form, which predominates in neutral aqueous solution, is not an electron-withdrawing group (EWG) but rather a donor or even neutral\(^{23,26}\) group; some photochemical decarboxylations were also reported.\(^{28}\) Thus, the promising photochemical properties of probe 6 convinced us to design the new PD-ABPP probes 7–11 (Figure 1B) functionalized by different EWGs in para-position of the benz(o)yl chain and an additional reporter group (i.e., alkyne prone to be reactive with azides in the click reaction). Noteworthy is that the p-alkyne group can be considered both as the reporter group for the CuAAC reaction but also an EWG to favor the formation of an insertion product upon photoirradiation.\(^{29,30}\)

**Synthesis of Clickable 3-Benz(o)ylmenadiones as PD-ABPP Probes**

Each of the 3-benzylmenadiones alkyne derivatives 2, 4, 6, and 8 was synthesized using as a key step, the Friedel–Crafts acylation, recently described by our team.\(^{31}\) Using the electron-rich 1,4-dimethoxy-2-methylnaphthalene (i.e., aromatic nucleophile) and readily available synthetic benzoic acids (7a–10a) (i.e., acylating agent) as starting materials, this reaction variant allowed us to prepare the corresponding 2-benzoyl-1,4-dimethoxynaphthalene intermediates (7b–10b) in mild conditions. These are key intermediates, to achieve, in a few additional steps the desired chemicals probes described in paths A and B (Scheme 1). The four benzoyl-1,4-dimethoxy-2-methylnaphthalenes 7b–10b were obtained with 47%, 66%, 82% and 67% yield, respectively. It is worth mentioning that without the Friedel–Crafts reaction variant, the 3-benzoylmenadiones functionalized by an alkyne group could not have been produced easily (i.e., see the different synthetic pathways discussed previously).\(^{31}\)

A Sonogashira pallado-catalyzed coupling allowed the trimethylsilane (TMS)-protected alkyne insertion, starting from the iodinated aromatic compounds 7b–8b, to obtain efficiently intermediates 7c–8c. These were successively deprotected in 7d–8d, first with TBAF and then by cerium ammonium nitrate (CAN) to afford both desired alkylated 3-benzoylmenadiones 7–8 upon oxidative demethylation. For the synthesis of alkynes 9c and 10c, propargyl alcohol was first submitted to a nucleophilic aromatic substitution reaction on the electron-poor fluorinated aromatic intermediates 9b and 10b, leading to the targeted quinones 9 and 10 after oxidative demethylation with CAN.

ABPP probe 11 in the benzylmenadione series was synthesized according to path C in a five-step route starting from the Kochi–Anderson reaction.\(^{12}\) (Scheme 1). First, 3-benzylmenadione 11a (80%) was produced by addition to menadione of a benzyl radical generated from 4-iodophenol-acetic acid by decarboxylation in the presence of silver salts’ catalysis and stoichiometric amounts of the Na\(_2\)S\(_2\)O\(_8\) oxidant. Owing to the incompatibility of the methyl group of 11a in basic medium, it was not possible to introduce the alkyne moiety directly on the quinone by palladium cross coupling reaction. Consequently, the benzylmenadione 11a was first reduced with SnCl\(_2\) in acid medium to the corresponding 2-(4-iodobenzyl)-3-methylnaphthalene-1,4-diol intermediate, which was protected (after a quick crystallization step under nitrogen) by methylation using dimethylsulfate to produce the 2-(4-iodobenzyl)-1,4-dimethoxy-3-methylnaphthalene intermediate 11b (56%). Then, the iodo derivative 11b was submitted to the Sonogashira pallado-coupling reaction, using ethynyl(trimethyl)silane in excess. This reaction successfully promoted the formation of the TMS-protected alkyne 11c in excellent yield (90%). The TMS group was removed from 11c by TBAF to obtain the free terminal alkyne 11d (97%), and the 1,4-quinone moiety was recovered by oxidative demethylation following addition of CAN in acetonitrile (ACN)/water mixture to obtain in good yield (66%) the targeted 3-benzyl-[4’-alkynyl]-menadione 11.

**Standardization of the UV Cross-Linking Parameters Using PD-ABPP Probes and nMet as Partner Model**

These PD-ABPP probes have been primarily designed to highlight both the binding sites and elucidate the protein interactome of PD in living cells. Additionally, considering the originally studied glutathione reductases from the pRBC unit, the reduction site of subsersive substrates is a matter of debate (discussed in ref.\(^{33}\)), even if menadione was observed to bind to various cavities of the crystal structure of the human enzyme.\(^{34}\) Before testing the cross-linking ability of these ABPP probes, we undertook the evaluation of their inhibition capabilities with the human GR in 1 mM GSSG reduction assays (in 2% ACN instead of 1% DMSO). We observed that the probes 7, 8, 9, and 10 behaved as potent GR inhibitors, with IC\(_{50}\) values of 0.60, 0.80, 0.58, and 0.85 \(\mu\)M, respectively; these values are in the same range as those previously observed for 6 and 12 (0.4 and 0.7 \(\mu\)M, respectively).\(^{17}\) Thus, functionalization did not decrease the inhibitor activity in comparison to effective 3-benzoylmenadiones.

Photoreactivity of the PD-ABPPs was evaluated under the same photoreaction conditions used for the model reaction between nMet and probe 6 (vide supra). We observed that among the five PD-ABPP, the photoreaction preferably occurred with probes 6, 7, and 9 bearing NO\(_2\) or alkyne (strong to moderate EWG) in para-position in accordance with yields of starting probe consumption and photoproduct formation, calculated from the \(^1\)H NMR spectra (Table 1,
Table 1. Photoirradiation of N-Acetyl Methionine Acid Methyl Ester (nMet) and Probes 6, 7, and 9

| probe | δ (ppm)/signal shape for each probe and photogenerated product from 1H NMR spectra | residual probe:formed products (ratio) |
|-------|---------------------------------------------------------------------------------|--------------------------------------|
| 6     | 8.07 (dd)/8.04 (dd)                                                             | 69:31                                |
| 7     | 7.60 (dd)/7.38 (dd)                                                             | 75:25                                |
| 9     | 7.51 (dd)/7.40 (dd)                                                             | 61:39                                |

“A Ratio of residual probe and formed coupling products is based on the integration of the corresponding signal in the 1H NMR spectra of the crude reaction mixtures.

Figures S2–S4). For probes 8, 10, and benzophenone, the 1H NMR data did not allow the calculation of the yields of probe consumption or formed products because the signals were either too small or combined with other signals. The CF3 group is known to be a strong EWG group, such as -NO2 and even more than an alkyne group27 but the mesomeric effect, inherent to both of the latter groups, can stabilize more efficiently a ketyl radical generated by photoreduction, than the inductive effect of a -CF3 group. Additionally, we determined the absorption spectrophotometric characteristics (εmax and λmax) of all PD-ABPP probes 7–11 along with probe 6 (Figure S5; at the photoirradiation λ of 350 nm, all the PD-ABPP probes are characterized by weak n–π* transitions). On the basis of these data, we can conclude that benzoylmenadione probes are photoreactive per se and can be used to covalently trap targets in an ABPP approach.

The (Pro-)ABPP Benzylmenadione Probe 11 Generates the Benzoylmenadione 7 upon Photoreduction and Then Oxidation

Starting from the prior observation of the generation of trace amounts of oxidized and reduced benzoylmenadione species upon photoreirradiation of the parent 3-benzylmenadione, we investigated the reaction to produce the most efficient photoreactive probe 7 from the parent prodrg 3-benzylmenadione 11. Since it is well-known that the dihydroenaphthoquinone is favorably generated via the semiquinone by photoreduction of the naphthoquinone in isopropanol,36 we submitted the benzylmenadione probe 11 to UV-irradiation in this solvent under bubbling of oxygen. After 72 h of irradiation, the formation of the corresponding 3-benzoylmenadione was clearly observed, as attested, for example, by monitoring the deshielding of the alkyne proton seen in the NMR spectra (Figures 3, S6, for the full-scale 1H NMR spectrum). The same reaction in a 1:1 mixture of ACN and water only afforded traces of the species oxidized at the benzylic position. To overcome the poor solubility of probe 11 in both systems, a 1:1 mixture of dichloromethane and isopropanol was selected, and these conditions were shown to be optimal with a full conversion of the initial 3-benzylmenadione 11 to the 3-benzoylmenadione 7 upon photoirradiation. The aerobic benzylic oxidation mechanism likely involves intermediate hydroperoxidation.36 However, the low quantity of 3-benzoylmenadione generated during the photoirradiation of 3-benzylmenadione in an ACN/water mixture prevented any detailed study on the optimization of the conditions for effective photoalkylation followed by the click reaction. This condition is not physiological, but we could demonstrate herein that naphthoquinone reduction by a flavoenzyme (e.g., both GRs) can be substituted by UV-photoactivation in the presence of oxygen. Notably, the benzyl oxidation might occur in living cells during enzymic catalysis with flavoenzymes in a hydrophobic environment like membrane lipid bilayers or a hydrophobic protein core. For this reason, the following study was carried out using 3-benzoylmenadione-based ABPP probes (instead of 3-benzylmenadione-based ABPP probes) to optimize the methodology and strengthen the ABPP approach.

Characterization of Clickability Properties of the ABPP Probes Using Azide Models

The click (CuAAC) reaction is crucial for the analysis of complex proteomes because it allows the grafting of a pulldown-tag to the cross-link adducts. Subsequent adduct enrichment through the affinity purification enhances correct peptide identification during MS analysis. The ABPP probes 7–10 were predicted to have different click reaction reactivity depending on the position of the alkyne on the phenyl ring and the length of the linker connecting them (via O–CH2 or directly attached). To assess the influence of both factors on the ABPP properties and select the best probe in model click reactions, we first evaluated the PD-ABPP reactivity with the commercially available and fluorescent rhodamine azide (RA) (Figure S7C). RA was used to develop and improve the reaction conditions by varying Cu(I) ligands (TBTA, THPTA, or BCDA) and/or the reductants (NaAsc and TCEP) that are essential for the efficiency of the CuAAC reaction (Figure S7A,B). The yields of the CuAAC reactions were determined by LC-MS analysis (Figures S8–S15).

During the development of an optimized protocol for the click reaction, we identified several factors, which surprisingly have greater than anticipated influence on the effectiveness of the click reaction with ABPP probes 7–10. Although well-known, the influence of these factors has not been sufficiently emphasized and described in the literature and has led us to
carry out a detailed fundamental investigation of each of the partners of the CuAAC reaction (vide infra). Additional observations, troubleshooting and click reaction optimization steps are described in the Supporting Information.

To improve the performance of the CuAAC reaction, we applied the generally used CuSO₄-THPTA-TCEP (copper source-ligand-reductant) trio in a 1:1:1 ratio. According to the yield of the optimized click reaction (Figure S10B), the sequence of probe efficiency (i.e., 2 h reaction) was determined as follows: probe 7 with -p-alkyne (58.8% yield) > probe 9 with -p-NO₂ and m-O-CH₂-alkyne (12.8% yield) > probe 10 with -p-CF₃ and m-alkyne (2.9% yield) > probe 8 with -p-CF₃ and m-alkyne (2.2% yield). The CuAAC reaction efficiency can be directly correlated with the probe structure.
and the resulting three factors: steric effects around the alkyne group, aqueous solubility of the probe and EWG properties of the aryl side groups (See Supporting Information, section “Click reaction optimization and troubleshooting”).

Additionally, in a click reaction with probe 7, we compared the generally used THPTA ligand with another Cu(I) ligand, the bathocuproinedisulfonic acid (BCDA) 37 in various conditions of the (copper source−ligand−reductant) trio both in water and in PBS buffer (Tables S1 and S2).

With this optimization study, we could conclude that phosphate ions can inhibit the CuAAC reaction and that this problem can be solved by lowering the phosphate buffer concentration and increasing copper/ligand ratio with respect to TCEP (Figure 4). Under these newly designed experimental conditions, we demonstrated that probe 7 can be clicked with an efficiency as high as in water without increasing concentrations of the reductant. BCDA is fully compatible with this click reaction conditions in PBS buffer (Table S2, R28−R30). Furthermore, it is preferred over THPTA in oxygen-free conditions.

Finally, we analyzed the click reaction of probe 7 with biotin-azide (BA), which is used to enrich tagged adducts by interaction with streptavidin. Despite changing the cosolvent of the reaction medium from DMF to ACN, the Cu(I) cycloaddition of BA had a similar pattern in triazole formation efficiency as RA (R32−39 vs R40−48; Tables S3 and S4). Thus, we conclude that our optimized click conditions are also compatible with an efficient labeling of alkynes with the biotin tag.

**Using Peptide as a Model for Photoreaction**

Based on nMet-PD-ABPP cross-linking data, we chose probes 7 and 9 to further explore the cross-linking ability of the ABPPs toward a peptide model. Additionally, this allowed us to determine the peptide adduct behavior (mass shift, fragmentation) during MS analysis, which is a crucial parameter to facilitate detection in proteomic analysis. The GSH and P52C peptides were chosen as models for cross-linking. GSH was selected as a model peptide because of its commercial availability, simple structure, high solubility in water, and relevance in the context of our model approach. P52C is a 16 amino-acid-peptide covering the pseudoactive site of the trypanothione disulfide oxidoreductase from T. cruzi.39

To assess the photoactivation of ABPPs in more physiological conditions, cross-linking with GSH or P52C was performed in a water:organic solvent mixture (H2O:ACN, 1:1, v/v). These conditions were different from those used for the cross-linking conditions with nMet (vide supra) where pure ACN solvent was used. However, we observed that the probe solubility is significantly limited in aqueous ACN solutions (Figure S16). The observed solubility properties of the probes followed this sequence (from the less to the most soluble): probe 10 < probe 8 ≈ probe 7 < probe 9. Since probe 9 was the most water-soluble ABPP probe and displayed the highest cross-linking efficiency with nMet, we used it as a binding partner for GSH. We found that although reagent concentrations were lowered (from mM to μM), we were still able to identify a significant fraction of GSH/GSSG-probe adducts after overnight photoirradiation (Figure 5A,B).

MS/MS analysis of the most prominent product (681.16 Da, retention time (RT) = 33.5 min) revealed that this adduct has apparently lost two hydrogens (expected M-2H) compared with the initially expected mass of the photoalkylated peptide (Scheme 2, pathway 1). However, MS fragmentation of this adduct revealed no further alteration from predicted peptide fragmentation patterns (Figure 5C). Similarly, cross-link adducts have been detected for P52C (Figure S17).

A more rational explanation for the apparent loss of two hydrogen atoms of the observed probe 9-GSH adduct could stem from the second pathway (Scheme 2) upon photoreduction of probe 9 in the presence of GSH. After photoreduction, an intramolecular process, much faster because it is entropically favored, leads to the benzoxanthone

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**Scheme 2. Chemical Analysis of the Insertion Products upon Photoirradiation of the ABPP Probe 9 with Glutathione (GSH) by Mass Spectrometry**

Two pathways of photoreactivity of the benzoylmenadione were expressed through the formation of two insertion products (blue box) with nucleophilic partners.
Several studies have clearly exemplified the photoreduction of quinones and subsequent intramolecular cyclization of a phenoxy radical, which effectively occurred in the presence of a H-donor. It is important to note that in these experiments, GSH can act as both reductant and H-donor.

Phenolate radical in position C-4 of the diradicaloid reduced intermediate promotes the intramolecular oxidative phenolic coupling. The methyl group in ortho to the free phenolate radical of the resulting benzoxanthone possesses a very labile -H, which releases a benzoxanthone-derived enone owing to the favored energetically structure. The electrophilic enone undergoes Michael addition with GSH leading to the benzoxanthone adduct (theoretical m/z = 681.16) in perfect agreement with the observed mass peak at m/z = 681.16. It is worth highlighting that the initial photoradiation of probe 6...
with nMet did show an additional mass peak \( m/z = 524.1 \), albeit with lower intensity, in the FD-MS spectrum (Figure 2A), attesting to the expression of two pathways occurring in the photochemical reaction.

Indeed, additional MS/MS analysis of the GSH adduct revealed that the generated probe fragment is benzoxanthone and that it was bound to the peptides at the sulfur atom of the cysteine residue (Figures 6C, S18). Consequently, a major formed probe species with the retention time of 40.2 min and \( m/z = 376.08 \) (identical to the probe 9 mass) found after photoirradiation was identified as the benzoxanthone (Figure 6B,C). This compound was not detected in the nonirradiated control (Figure S19A) or after 10 min of irradiation (Figure 6A), suggesting that prolonged photoreduction time is necessary to generate the cyclization product. Additionally, the newly found species underwent deprotonation forming the predicted and reactive enone of pathway 2 \( m/z = 374.07 \) (Figures S20, S21E). Incubation of synthesized PDO-BX with GSH confirmed the BX reactivity toward free thiol of GSH (Figures S22A, S22B, S23).

Interestingly, although no benzoxanthone is formed after 10 min of UV-irradiation of PD metabolite PDOox or probe 9, with GSH, the reactions also gave rise to adducts missing two hydrogen atoms (Figures S2A, S22C). MS/MS analysis identified this compound as a 2-(S-glutathionyl-substituted-methyl)-3-(benzoyl)-1,4-naphthoquinone (shortened as 2-(GS-methyl)-PDO or 2-(GS-methyl)-probe 9) (Figures S24A, S25). Surprisingly, the 2-(SG-methyl)-9 is not present upon overnight irradiation of probe 9 and GSH, suggesting that the species is an intermediate formed in the synthesis of 9-BX-SG, according to pathway 3 (Scheme 3).

To further support our findings on the occurrence of pathways 2 and 3 occurrence, we substituted GSH in the reaction with another nucleophilic agent with a thiol group—thiophenol. LC-MS showed that already after 10 min of irradiation of PDO or probe 9, benzoxanthones as well as adducts lacking two hydrogens were formed (Figures S26, S27).

However, the suggested pathways are not mutually exclusive as a more careful LC-MS/MS analysis of the probe 9 reaction mixtures with GSH or thiophenol revealed that formation of benzophenone-like adducts occurred as well (Figures 6B, S24B, S26B, S28). Furthermore, in photoreactions, the nitro group from probe 9 was photoreduced to an amine,\(^3\) which has given rise to amine-substituted benzophenone adducts and -(SG-methyl)-9 adducts (Figures 6B, S29, S30).

With that, we demonstrated that probe 9 is able to efficiently cross-link to a peptide and that the corresponding peptide-ABPP adducts can be detected by MS analysis.

Importantly, three labeling pathways were evidenced to occur in the photoirradiation experiments involving the metabolite PDOox or probe 9 and GSH, as depicted in Schemes 2 and 3. Using the LC-MS/MS approach, we were able to detect the main intermediates and products of the

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\[ \text{Scheme 3. Mechanism of Formation of Both Observed Insertion Products (Blue Box) via Pathway 3 upon Photoirradiation of the ABPP Probe 9 with Glutathione}^{\text{a}} \]

\[ \text{The structure of the intermediate 2-(SG-methyl)-probe 9 adduct, formed after 10 min-irradiation, was deduced by ESI-MS/MS mass spectrometry.} \]

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pathways: the probe, the benzophenone-like adduct, the 2-(SG-methyl)-probe adduct, the cyclized probe-BX, the probe-BX-derived enone, and the probe-BX insertion adduct.

**Probe Cycling with Glutathione Reductase Generates Benzoxanthone**

As for photoreduction, the benzoxanthone formation has been postulated to occur during several cycles of enzymatic (GR) 1e-reduction of PDOox (Figure 1A). However, the metabolite was only indirectly detected by electrochemical measurements of PDO derivatives due to its minor amount. To prove definitively that PDO-BX is generated by continuous redox-cycling of the drug under hGR catalysis, we analyzed such reaction by LC-MS/MS after 6 h of regular addition of NADPH. The BX-derived enone could be found in reactions in open air (Figures S21B, S21C) but not in the deoxidized control where redox-cycling was not possible due to the absence of oxidants like oxygen (Figure S21A). This clearly demonstrates that PDO-BX is indeed a product of PD metabolite redox-cycling (Figure 1A).

In addition, we investigated similarities in the BX formation during the redox-cycling processes during photoreduction and GR catalysis. For this, we irradiated hGR with probe 9 in oxygen-free conditions. Interestingly, despite the lack of oxygen, we were able to obtain 9-BX from probe 9 after 10 min of UV-irradiation with hGR (Figure S21D), although this was not possible in a comparable period of time when GSH was acting as a nucleophile. This demonstrates that the presence of the enzyme is enough to accelerate light-induced formation of 9-BX. Indeed, the UV-photoreduction process can mimic the reduction of naphthoquinone by NADPH-reduced enzyme in this pathway, indicating that both processes might share similarities. Generation of BX from PDOox or probe 9 was also possible in the presence of thiophenol after 10 min of photoirradiation. However, the cysteine thiol group in GSH only led to minor formation of PDO-BX even after overnight UV-irradiation (Figure S22 compared to S26). The results obtained with hGR upon irradiation imply that the protein cysteines might be more reactive than GSH. Alternatively, the entropic interaction between the naphthoquinone and the enzyme might play a mutual influence on each other upon transferring electrons and kinetically favor pathways 2−3 following probe binding to a cavity where the molecular environment favors BX formation.

**Using Glutathione Reductase as a Model for Photoreaction**

To test the ability of the probes to interact with protein targets and to understand the exact probe binding/reduction site in GR structure we studied the cross-linking pattern of an ABPP
probe with GR upon photoirradiation. Enzymatic inhibition assays proved that the ABPP probes series (e.g., probes 9 and 6) with a EWG (i.e., NO2) in para-position to the benzoyl moiety possess the highest inhibitory potency toward hGR among all probes. Furthermore, considering the better solubility and photoreactivity of probe 9, we selected it as the best ABPP probe for studies with model proteins hGR and PfGR.

Mass spectrometry analysis confirmed probe 9-hGR or -PfGR adduct formation. Similarly to GSH cross-linking, the adduct m/z suggested the formation of GR-probe products such as benzophenone-type adduct (+376 Da) and 9-BX or 2-(S-methyl)-9 (+374 Da) adducts as well as species with reduced nitro group (346.11 and 344.10 Da)(Figure S31).

Importantly no significant cross-linked peptides (Mascot score >30) have been identified during MS analysis of UV-irradiated negative control BSA incubated with 9 and the UV-irradiated GRs alone.

Interestingly, the cross-linking sites associated to 9-BX or 2-(S-methyl)-9 were found in sites distinct from the active cysteines. Alkylation of GR was pinpointed at Lys residues whose intrinsically nucleophilic ε-amines confer important roles. Beyond cysteines, lysines represent a source for covalent probe development, and several studies on epigenetics in vivo have started to identify and map the Lys ligand ability of the human proteome.45 We can assume that the highly electrophilic BX-derived enone might be attacked by the intrinsically nucleophilic lysine ε-amine (Scheme 3). Furthermore, Lys

Figure 8. Probe 9 cross-links to PfGR. (A) Positions of cross-linked amino acids (K415 — blue, V6 — pink) to probe 9 in the PfGR structure. The substrate binding cleft leading to the catalytic disulfide bridge and interspace cavity opening is indicated by the orange triangles. (B) Images picturing the distance between cross-linked K397 in hGR (left panels), K415 in PfGR (right panels), and the interspace cavity opening. Upper panel — amino acid positions on protein chain. Lower panels — surface density of cavity opening. Blue — cross-linked lysine; Green and violet — amino acids of the cavity opening.
cross-linking would explain why this amino acid is misincised after trypsin digestion in a large portion of identified peptide adducts (Figure S31). Similar cleavage blocking has been reported in MS studies when Lys is targeted by electrophiles and supports our correct identification of the adducts.45

Analysis by MS/MS of the probe 9-cross-linked peptide adducts identified three hGR peptides with high confidence (Figures 7A, S31A). The identification of these adducts several times confirms that the probe 9 can be used as a photocative ABPP probe to bind to target proteins. The involved binding site between K255 and T257 on peptide E253-R272 is located on the exposed surface of the protein, away from the protein catalytic centers, in this apparently nonfunctional region. However, the two other identified binding sites lie in known functionally related regions.

The cross-linking position (K397) of peptide Y393−K416 does not reside in known binding pockets of hGR (Figure 7A,B). Interestingly, the binding locus correlates to previous enzyme kinetic analyses performed with 3-benzoylmenadiones exhibiting uncompetitive GR inhibition.47 Indeed, it is localized in the interface domain, known to mediate the dimerization of the protein. The domain is crucial for GR activity as the enzyme is not active as a monomer. Additionally, in the interface domain, at the 2-fold symmetry axis of the homodimeric protein, a cavity is present, which was reported to bind numerous GR inhibitors, such as 2-methyl-1,4-NQ (menadione),34 6-hydroxy-3-oxo-3H-xanthene-9-propionic acid,46 a series of 10-arylisoalloxazines,47 and S-(2,4-dinitrophenyl) glutathione.48 The cavity does not have a direct connection with the NADPH or FAD binding sites. It is linked to the external surface of the protein and to the GSGG binding site by two pairs of short channels. The channel pair openings, which are located in the catalytic center, emerge at the bottom of the V-shaped catalytic crevices in close proximity to the redox-active disulfide bridges. Significantly, in total, 41 amino acids of both subunits participate in the GSGG binding site in hGR.36 It was suggested that compounds docked in the cavity37 could either trigger structural changes disturbing the dimer stability leading to decrease of enzyme activity or interfere with the redox potential of the flavin.

Interestingly, the amino acids of the channel opening (403-FTPMYH-408) are present in the identified peptide where K399 was alkylated by probe 9, however, separated by a distance of 19−25 Å from the identified cross-link site and 9.5−19 Å from the active Cys in the GSGG binding site. It is possible that the original interaction site does not overlay with the alkylation site especially considering the reactivity of Lys toward BX. In line, when probe 9 was cross-linked with PfGR, instead of hGR, the peptide that was identified with the highest confidence (I411−K431)(Figure S31B) is the exact homologous region of peptide Y393−K416 in hGR (Figure S31A). Moreover, compared with hGR, the suggested part cross-linked to the probe in this peptide (411-IFESKFT-417) contains a K415 to S402 substitution, which lies in proximity to the channel opening (Figure 8A,B). K415 was misincised during trypsin digestion indicating the exact probe modification site at this amino acid. Strikingly, the peptide sequence prior to the cavity is conserved between PfGR and hGR with a high degree of identity emphasizing the importance of this region (Figure 8).

The amine generated upon nitro group reduction in the 9-BX[NH2]-peptide S225−K247 adduct was localized on the free cysteine Cys234 lying in proximity to enzyme’s catalytic center where FAD is reduced by NADPH, especially very close to the ultraconserved tyrosine 197. Tyr197 serves as gate keeper of cofactor access to FAD because of its ability to flip and block NADPH positioning near FAD.50

Structural analysis shows that Cys234 is part of a sizable pocket that extends into it all the way from the surface near the side chain of Ala 241 (Figure 7C). The pocket lies in a relatively accessible region, which directly interacts with a network of water molecules. Since the pocket is partly lined with nonpolar side chains, this could allow the naphthoquinone to displace the water molecules and gain access to Cys234. Trapping of the probe could additionally be bolstered by intrinsic reactivity of the 9-BX toward cysteine. In fact, Cys234 might have promoted the formation of 9-BX via pathways 2 or 3 (Schemes 2 and 3). When testing the orthologous Plasmodium enzyme PfGR, cross-linking with probe 9 did not occur at the homologous peptide, probably because of the lack of cysteine in this region in comparison to the equivalent in hGR. Of note, the pocket does not appear to have direct access to Tyr197 as it is additionally blocked by Val200 or Cys234 itself. Nevertheless, Cys234 and especially the surrounding cavity represent an interesting target for future PD MoA investigation. A clearer picture of its importance should be studied in the future by generating hGR mutants for enzyme kinetics and drug binding/reduction evaluation.

Photolabeling of hGR and Pull-Down of Labeled Protein Adducts

Having established the cross-linking and click conditions for the probes as well as selection of the most efficient ones for ABPP, we tested their labeling capability of proteins on hGR. Cross-linking with 9 and subsequent click reaction allowed for efficient RA attachment on hGR (Figure S32). Similarly, tagging with BA after cross-linking with 7 or 9 followed by pull-down with avidine of labeled adducts proves the ability of the probes to target and isolate proteins (Figure 9). Both tagging reactions of probe 9 with RA and BA are competitive toward nonclickable drug analogues (6 in RA and PDO in BA labeling), demonstrating the specificity of the labeling. Altogether, we evidenced that both the cross-linking and click reaction of our probes can be combined for the ABPP strategy.

![Figure 9](https://doi.org/10.1021/jacsau.1c00025) Pull-down of hGR labeled with ABPP probes 7 and 9 and clicked with avidin tag. SDS-PAGE gel stained with Coomassie is pictured. For each reaction, 2% of the reaction before pull-down and 50% of the elution after avidin binding were loaded on the gel. hGR is localized at the height of the 55 kDa marker band. M − marker.
formation of a PDO–hematin complex by binding titrations.\textsuperscript{20} Having demonstrated the electrophilic nature of the BX 4-derived enone, we used a collision-induced dissociation tandem mass spectrometry (CID-MS) methodology to characterize the complex in detail.\textsuperscript{51} In particular, two intense PDO-BX\textsuperscript{−}heme adducts at \( m/z = 960.2 \) ([heme+PDO]\textsuperscript{−}) and \( m/z = 975.3 \) (formally [Heme+BX 4•H+O]\textsuperscript{+}) were clearly detected (Figure 10A). The first BX 4•-heme adduct at \( m/z = 960.2 \) corresponds to a \( \pi-\pi \) complex. This feature is assessed by the weak dissociation voltage \( DV_{50} \) (150 V) and the absence of a residual complex at high fragmentor voltage (Figure 10B).

In addition, the \( \pi \)-stacking in the PDO–hematin complex might be strengthened by additional interactions such as hydrogen bonding between the propionate of the heme side chain\textsuperscript{52} and the phenol in PDO-BX and Fe\textsuperscript{III} axial coordination of the BX carbonyl unit (Figure 10C, species A). It is noteworthy that PDO-BX can be oxidized (\( E_{pa} = -50 \text{ mV} \) and \( -208 \text{ mV} \)) through its phenolic moiety and might undergo an exchange of electron(s) with the tightly bound Fe\textsuperscript{III} heme. The intramolecular PDO → Fe\textsuperscript{III} heme 1e- transfer is favored by hydrogen bonding\textsuperscript{52} and generates a carbon radical at the BX core (Figure 10C, species B).

The concomitantly formed Fe\textsuperscript{II} heme from species B binds O\textsubscript{2}, and the resulting species C (Figure 10C) might be attacked by the nucleophilic \( \alpha \)-keto carbon radical\textsuperscript{53} of the tightly bound BX, ultimately leading to hematin meso-alkylation\textsuperscript{54} by PDO-BX, as suggested from the CID-ESI-MS experiments (second adduct at \( m/z = 975.3 \)). It is noteworthy that the \( DV_{50} \) value of this second PDO–heme adduct is markedly increased (+32 V), and a significant amount of the complex is observed at high fragmentor voltage, which is therefore indicative of a very stable heme adduct (\( \sim 5\% \) for PDO–heme adduct and \( \sim 7\% \) for the antimalarial drug amodiaquine (AQ)–heme adduct used as reported reference). In this experiment (Figure 10B), the antimalarial chloroquine (CQ), known to be a reversible heme binder did not show a residual covalent adduct at high fragmentor voltage. Similarly, when probe \textsuperscript{9} was UV-irradiated with GSH, the formed benzoxanthone was demonstrated to be reactive toward heme, when added to the reaction, leading to the generation of the adduct 9-BX–heme and its hydrated version (Figure S33). Together with the

Figure 10. ESI-MS and CID-MS analysis of PDO-BX 4-heme complexes. (A) ESI mass spectrum (exit potential: 120 V) of a 1:1 mixture of 50 \( \mu \)M heme and 50 \( \mu \)M PDO-BX 4 in H\textsubscript{2}O/CH\textsubscript{3}CN (5/95) – 1% formic acid. (B) Stability responses of the BX 4-heme (at \( m/z = 960.2 \) and at 975.3), AQ-heme (at \( m/z = 971.3 \)) and CQ-heme (at \( m/z = 935.4 \)) complexes obtained by CID-MS experiments. ESI-MS\textsuperscript{−}; 120 V < fragmentor< 400 V with 20 V increments. (C) Proposed molecular structure of iron\textsuperscript{III}-hematin species alkylated by the BX 4 and comparison between simulated and observed mass signatures of species A and D.
CID-MS experiments, this proves that the previously observed PDO-BX—heme complex is covalently linked to heme through the reactive enone alkylation. The structural signature of this alkylated hematin product is tentatively proposed in species D (Figure 10C), following reaction of the quinone methide radical at the meso-position of the tetrapyrrole and release of a water molecule, as already demonstrated for artemisinin.

This suggested that the heme alkylation product has to be regarded as the result of the formation of a key carbon radical generated from a redox-active agent in redox-driven bioactivation processes and a relevant reaction to the MoA occurring in the parasite in vivo.

Such contribution needs more detailed investigations to understand the MoA of the redox-active lead animalarial PD. Interestingly, the data obtained with PDO-BX are reminiscent of the hypothesized formation of xanthones to explain the potentiation of antimalarial activities of polyhydroxylated benzophenone derivatives tested in the presence of Fenton catalysts upon catalysis of redox-active metals such as Fe3+.

In the present study, upon oxidative phenolic coupling of PDO_red, BX releases a powerful electrophile that can be attacked by the nucleophilic species present in the reaction, GSH, the terminal ε-amine group of lysine-like K397 in hGR, or heme. Evaluation of the Antimalarial Properties of PD-ABPP

To validate the applicability of ABPPs in parasites, we then evaluated the antimalarial activity of the five newly synthesized 3-benz(o)ylmenadione-based ABPP probes (7–11, Table 2).

Table 2. IC50 Values for 3-Benz(o)ylmenadione Derivatives Determined from Growth Inhibition Assays with Highly Synchronized P. falciparum Strain Dd2

| series            | compound | P. falciparum Dd2<sup>a</sup> (nM) | hMRC-S<sup>b</sup> (μM)<sup>c</sup> |
|-------------------|----------|----------------------------------|-----------------------------------|
| 3-benzoylmenadiones | 6        | 513 ± 287                        | 24.0                              |
|                   | 7        | 1806 ± 302                       | 20.5                              |
|                   | 8        | 2993 ± 750                       | 25.8                              |
|                   | 9        | 417 ± 222                        | 42.2                              |
|                   | 10       | >5000                            | 29.4                              |
| 3-benzylmenadiones | 11       | 49 ± 15                          | >64.0                             |
| plasmodione       |          | 20 ± 5                           | >32.0<sup>d</sup>                 |

<sup>a</sup>Three independent experiments with the SYBR green assay (mean ± SD).<sup>b</sup>The P. falciparum Dd2 strain is sensitive to DHA (IC50 DHA = 0.7 ± 0.2), to methylene blue (IC50 MB = 7 ± 0.3), and resistant to chloroquine (IC50 CQ = 189 ± 12).<sup>c</sup>Toxicity values against human fibroblasts hMRC-S were determined by using reported protocols.<sup>d</sup>Value from ref 17.

The potent antimalarial activities of plasmodione 1, its nitro analogue 5 and the PDO-BX 4 were already reported.<sup>17,20</sup> As previously observed and despite being the likely key metabolites of 3-benzylmenadiones, the 3-benzyolmenadiones do not display a high antimalarial activity, with an IC50 of ca. 10–50-fold higher than the corresponding 3-benzylmenadiones.<sup>17,21</sup> This may be explained by the very poor internalization of 3-benzylmenadione metabolites in pRBCs when given externally. Indeed, similar to the 3-benzyolmenadione metabolite PDO, probes 6–10 (Figure 1) are more polar and planar than the 3-benzylmenadiones (PD, probes 5 and 11), likely reducing their ability to be internalized in parasites, and in accordance with the same observation in the yeast model.<sup>57</sup> To act as the key active principle of the prodrug PD, the metabolite has thus to be generated in situ in the parasite before it can efficiently cycle with NAD(P)H-dependent reductases. With respect to the ABPP properties studied in the click reaction and under photoirradiation, we observed that probes 7 and 9 are the most efficient probes to be used in photolabeling of plasmodione targets.

This result has motivated the synthesis of a first PD-ABPP, probe 11, in the 3-benzylmenadione series. Probe 11 (Figure 1) displayed an IC50 value comparable to that of PD regardless of substitution of the CF3 function at the para-position by an alkene group. On the basis of our studies on the photo-reactivity and clickability of 3-benz(o)ylmenadione-based ABPP probes, probe 11 is selected for future PD interactome analysis as the most efficient prodrug in killing parasites with a similar IC50 value as the value of PD and because it is expected to generate the most photoreactive probe 7 upon bioactivation in living Plasmodium parasites.

CONCLUSIONS

Herein, we have presented the design and the synthesis of specific (pro-)PD-ABPP probes based on the postulated MoA of the antimalarial prodrug PD. We have studied the influence of different EWGs in the 3-benzylmenadione series on the photo-reactivity of the ABPPs as well as the probes’ “clickability” properties. This allowed us to select probes 7 and 9 as the most effective tools for the ABPP approach. Optimization of the ABPP methodology (e.g., click in PBS with Cu<sup>2+</sup>:BCDA:TCEP (5:1:1)) has been successfully demonstrated by hGR photolabeling with probe 7 or 9 and subsequent pull-down of labeled protein adducts. Interestingly, labeling of different nucleophilic amino acids in distinct regions of hGR and PfGR will open new directions to study GR mutants of these different residues in the context of drug development. Finally, using UV-photoirradiation, we provide evidence that (pro-)ABPP probe 11 can indeed be photooxidized in 3-benzylmenadione-derived ABPP probe 7. While these conditions are not physiological, this result further supports the current model for PD activation and MoA. Additionally, by correlating the efficiency of (pro-)ABPP with their antimalarial activity, we concluded that the (pro-)PD-ABPP probe 11 should be the most effective one to be used in parasite cultures to identify the PD interactome by proteomics analysis. Identifying PD targets will be essential to further optimize the properties of this compound series and to design more active and target-specific derivatives.

Beyond the investigated plasmodione-based tools, we believe that the concept of (pro-)ABPP can be further expanded as a generalizable and emerging strategy to investigate redox-active drugs with various biological properties (e.g., anticancer, antibiotic, antiviral, antiparasitic) to identify new protein targets, while noting that other warheads than menadione are also possible.

Linking drug effects from complex reactions to killing processes/targets in parasites is one of the most challenging steps when studying the MoA of any drug. When the latter is a redox-activable prodrug it is even more difficult because a wide range of metabolites can be generated in trace amounts during redox bioactivation. One solution dealing with this quantity limit is the development ABPP probes based on already known drug metabolites, keeping in mind that the method addresses a common limitation of exogenously added probe, which does not necessarily mimic the endogenous generation of an active
principle from a prodrug in the target living cell. Therefore, to circumvent this limitation, we designed a series of photo-reactive prodrugs, which upon UV irradiation generate the same drug metabolites as those formed through redox bioactivation, and we put this concept into practice using the antimalarial agent plasmodioline. Besides this applied and deep investigation about an antiplosomal early lead agent, we believe that understanding the various parameters influencing the CuAAC reaction, for example, Cu(I) preincubation reaction with reductant and ligand, organic solvent, reductant and ligand ratios, pH, buffer dilution, salts, counter-ions, is essential for the success of the CuAAC strategy. The interplay between these elusive factors is key in the setting up of the optimization of the CuAAC reaction, as demonstrated by our findings.

### MATERIAL AND METHODS

#### UV-Irradiation

Reactions were irradiated either with a 365 nm light generated by a UV monochromator of 1000 W intensity for 8 to 10 min or with a 350 nm light generated by eight RPR-3500A lamps of 200 W with a Rayonet photochemical reactor overnight at a distance of 3 cm from the light source.

#### Irradiation Experiments for Photobenzylic Oxidation of the (Pro-)ABPP Benzylmenadione Probe 11 to Benzylobenzaldehyde 7

First, 50 mg of 2-(4-ethynylbenzyl)-3-methylnaphthalene-1,4-dione 11 and 2 mL of the appropriate solvent were added in a tube. The mixture was agitated and bubbled with oxygen during 30 min. Then, under a positive pressure of oxygen, the tube was placed in a Rayonet photochemical reactor and irradiated at 350 nm for 72 h. The resulting mixture was extracted with dichloromethane if necessary, and the solvent was removed under reduced pressure. The reaction crude was directly analyzed by NMR spectroscopy.

#### Irradiation Experiments for Model Photoreaction

The photochemical reaction of N-acetyl-methionine methyl ester (N-Ac-Met-OMe, shortened as nMet) with benzophenone and benz(o) menadione 7 occurred in a pyrex tube (filter for hν < 300 nm) at a final concentration of 0.1 M at a final concentration of 0.1 M in ACN. The reaction was irradiated at 5 °C for 24 h in a Rayonet reactor (at 350 nm). Finally, the reaction was analyzed by field-desorption mass spectrometry (FD-MS), as seen in Figure 2. FD-MS of the reactions was performed at Heidelberg University according to the published protocol.58 Protein bands were cut out and subjected to trypsin metrically with a Cary 50 absorption spectrophotometer by adding a 6 μM/nMet sample at regular 2 h-intervals for the next day the solution was analyzed by HPLC-MS.

For heme alkylation by 9-BX, 1.5 mM heme solution in PBS was added to the reaction mixture containing GSH and probe 9 at completion. The resulting mixture was incubated for 2 h and analyzed by HPLC-MS. Hematin was generated from heme (Sigma) by basifying the heme solution with 2 M NaOH.

In the study about cross-linked adduct formation, protein photodissociation was performed with a mixture of 6 μM hGR or 5 μM P4GR (or BSA - used as negative control) with 6 μM or 5 μM probe 9, respectively, in 200 μL of 10 mM PBS buffer at pH 6.9 with 2% ACN. Probe solubility in 2% ACN was assessed spectrophotometrically with a Cary 50 absorption spectrophotometer by monitoring absorbance kinetics of decreasing ABPP probe concentrations, starting with 20 μM (Figure S16).

In reactions with hGR and the probe, 52.5 μM NADPH was added to initiate the redox-cycling. The reaction mixture was deoxygenized by seven alternative vacuum and Ar flux cycles (10 s each) in an anaerobic cuvette. Subsequently, the mixture was photoirradiated for 8 min, and afterward, 100 μL of 3X Laemmli buffer was added. The next day, the samples were separated on 10% SDS-Page gels, stained with Coomassie solution, and destained according to published protocol.69 Protein bands were cut out and subjected to trypsin digestion and HPLC-MS analysis.

#### Generation of 9-BX from ABPP Probe 9 upon hGR Redox-Cycling

In order to generate 9-BX, 40 μM of probe 9 was allowed to redox-cycle with hGR and 1.44 mM NADPH. Probe 9 stock solution was prepared in DMSO and added to the reaction mixture in the presence of 2% solvent final in 47 mM PBS buffer in 200 μL of total reaction volume. In the hemoglobin reduction assay, 80 μM methemoglobin was mixed additionally to the reaction. Redox-cycling was started by addition of a 6 μL-aliquot of 16 mM NADPH and 4 μM hGR. The same amount of NADPH was added at regular 2 h-intervals for the next 6 h. A control sample was deoxygenized by seven vacuum–argon cycles before first addition of the separately deoxygenized NADPH solution.

#### Generation of 9-BX from ABPP Probe 9 upon hGR Photoreduction

Probe 9 photoreduction in the presence of hGR was achieved by mixing 100 μM of the probe in 20% ACN with 4 μM hGR in 47 mM PBS buffer. Samples were deoxygenized by 7 alternative vacuum–argon cycles with longer argon cycles (15 s) than vacuum cycles (<6 s) to then analyzed by 1H NMR. The covalently cross-linked products were confirmed by TLC and NMR analysis.

### Inhibition Potency of ABPP Probes toward Human GR

The inhibition of hGR by ABPP probes 7–10 was evaluated by using a standard GR assay5 with 100 μM NADPH (Bioivet) and 1 mM GSSG (Serva) in GR buffer (47 mM potassium phosphate buffer, 200 mM KCl, 1 mM EDTA, pH 6.9) at 25 °C in a 1 mL-cuvette using the inhibitor concentrations ranging from 0 to 2 μM. The final concentration of ACN in the assay was 2%. Initial rates of NADPH oxidation by hGR (8 mU or 0.85 pmol) was monitored at 340 nm with a Cary 50 UV–vis absorption spectrophotometer.

### Photoreaction between the ABPP Probe 9 and Thiophenol, Peptides, and Proteins as Partners

For peptide cross-linking, stock solutions of GSH (Sigma) or P52C were prepared in 20 mM PBS at pH = 6.9. Stock solutions of PDO 2 or of probe 9 were prepared in pure ACN. For thiophenol cross-linking, the stock solution of the thiol was prepared in ACN. Then, the reaction was started by mixing 3 mM of peptide or thiophenol with 600 μM 9 or 100 μM 3 in a mixture of 1:1 ACN and 10 mM PBS in 200 μL volume. The reaction mixtures were deoxygenized by seven alternative cycles of vacuum and Ar flux in an anaerobic cuvette with longer argon cycles (15 s) than vacuum cycles (<6 s) to avoid ACN evaporation. The mixture was photoirradiated for 10 min or overnight (O/N), and the next day the solution was analyzed by HPLC-MS.

For heme alkylation by 9-BX, 1.5 mM heme solution in PBS was added to the reaction mixture containing GSH and probe 9 at completion. The resulting mixture was incubated for 2 h and analyzed by HPLC-MS. Hematin was generated from heme (Sigma) by basifying the heme solution with 2 M NaOH.

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In reactions with hGR and the probe, 52.5 μM NADPH was added to initiate the redox-cycling. The reaction mixture was deoxygenized by seven alternative vacuum and Ar flux cycles (10 s each) in an anaerobic cuvette. Subsequently, the mixture was photoirradiated for 8 min, and afterward, 100 μL of 3X Laemmli buffer was added. The next day, the samples were separated on 10% SDS-Page gels, stained with Coomassie solution, and destained according to published protocol.69 Protein bands were cut out and subjected to trypsin digestion and HPLC-MS analysis.

#### Generation of 9-BX from ABPP Probe 9 upon hGR Redox-Cycling

In order to generate 9-BX, 40 μM of probe 9 was allowed to redox-cycle with hGR and 1.44 mM NADPH. Probe 9 stock solution was prepared in DMSO and added to the reaction mixture in the presence of 2% solvent final in 47 mM PBS buffer in 200 μL of total reaction volume. In the hemoglobin reduction assay, 80 μM methemoglobin was mixed additionally to the reaction. Redox-cycling was started by addition of a 6 μL-aliquot of 16 mM NADPH and 4 μM hGR. The same amount of NADPH was added at regular 2 h-intervals for the next 6 h. A control sample was deoxygenized by seven vacuum–argon cycles before first addition of the separately deoxygenized NADPH solution.

#### Generation of 9-BX from ABPP Probe 9 upon hGR Photoreduction

Probe 9 photoreduction in the presence of hGR was achieved by mixing 100 μM of the probe in 20% ACN with 4 μM hGR in 47 mM PBS buffer. Samples were deoxygenized by 7 alternative vacuum–argon cycles with longer argon cycles (15 s) than vacuum cycles (<6 s) to
avoid ACN evaporation. The reaction was UV-irradiated for 10 min and the mixture was analyzed by HPLC-MS.

**Successive Cross-Linking and Click Reaction with hGR**

For hGR labeling 150 μL of 10 μM hGR in 12.5 mM PBS (potassium based) and 2% DMSO was UV irradiated in the presence of 10 μM probe 7 or 9 for 10 min. The reaction was beforehand deoxygenized by seven alternative cycles of vacuum and Ar flux. In competition assays 30 μM of probe 6 or PDO was added additionally. After a 10 min photoreduction, 3.3% DMF and 20 μM RA or 10 μM BA was added. The reaction was deoxygenized a second time, and 0.4% of deoxygenized SDS was added with a syringe. A click reaction was initiated by adding a 1:1:1 copper BCDA:CuSO4:TCEP 40 min-long preincubation mixture to a final concentration ratio of 132:660:132 μM, respectively, and finally volume of 200 μL. The reaction was incubated overnight at 30 °C. Reactions containing biotin azide (BA) were subjected to pull-down, whereas rhodamine azide (RA) reactions were mixed with 100 μL of 3x Laemmlı, heated at 60 °C, and separated by SDS-PAGE. Gel fluorescence was visualized by GelDoc EZ imager (BioRad) on a blue ray (excitation = 430–460 nm). The gel was stained by Coomassie staining after fluorescence analysis.

**Biotin Pulldown**

Biotin-protein adducts were pulled down by binding to avidin agarose beads (Pierce). Prior to use, the beads were washed five times with 1.5 mL of washing buffer (47 mM sodium based PBS, pH 6.9) and centrifuged at 5000 g for 1 min at RT. Unspecific sites on the avidin agarose beads were blocked by incubating the beads for 1.5 h at RT with 0.5 mM BSA. Overnight click reactions were diluted with 47 mM PBS with 0.3% SDS to 1 mL of volume and incubated for 1 h with beads at RT. The suspension was washed once with washing buffer + 0.05% Tween20 and once with washing buffer + 1% SDS, as well as once with washing buffer in-between, before, and after. Subsequently, the beads were centrifuged at 4500g for 1 min at RT. Bound proteins were eluted at 96 °C for 10 min with 80 μL of Laemmlı buffer. Eluted proteins were separated on 10% SDS-PAGE gel and stained with Coomassie stain.

**Protein Preparation for In-Gel Digestion**

The gel pieces were successively washed with 50 μL of 25 mM NH4HCO3 and 50 μL of ACN (three times) and dehydrated with 100 μL of ACN before reduction in the presence of 10 mM DTT in 25 mM NH4HCO3 (1h at 57 °C) and alkylation in the presence of 55 mM iodoacetamide in 25 mM NH4HCO3. For tryptic digestion, the gel pieces were resuspended in 2× volumes of trypsin (12.5 ng/μL; Promega V5111) freshly diluted in 25 mM NH4HCO3 and incubated overnight at 37 °C. The digested peptides were then extracted from the gel in a buffer containing 34.9% H2O, 65% ACN, and 0.1% HCOOH, and the excess of ACN was removed by evaporation and peptides analyzed by nanoLC-MS/MS.

**NanoLC-MS/MS Analysis**

Peptide digests analysis was performed on a nanoACQUITY Ultra-Performance-LC (Waters, Milford, MA, U.S.A.) coupled to a TripleTOF 5600+ mass spectrometer (Sciex, Framingham, U.S.A.). The samples were trapped on a 20 × 0.18 mm, 5 μm Symmetry C18 precolumn (Waters Corp.), and the peptides were separated on a nanoEase M/Z Peptide BEH C18 Column, 130 Å, 1.7 μm, 75 μm × 150 mm (Waters). The solvent system consisted of 0.1% formic acid in water (solvent A) and 0.1% formic acid in ACN (solvent B). Trapping was performed during 3 min at 5 μL/min with 99% of solvent A and 1% of solvent B. Elution was performed at a flow rate of 300 nL/min, using 1–40% gradient (solvent B) over 35 min at 60 °C followed by 65% (solvent B) over 5 min. The mass spectrometer was operated in positive mode, with the following settings: ion spray voltage floating (ISVF) 2300 V, curtain gas (CUR) 25 psi, interface heater temperature (IHT) 75 °C, ion source gas 1 (GS1) 2 psi, declustering potential (DP) 100 V. Information-dependent acquisition (IDA) mode was used with top 5 MS/MS scans. The MS scan had an accumulation time of 250 ms on m/z [400–1250] range and the MS/MS scans 100 ms m/z [150–1600] range in high sensitivity mode. Switching criteria were set to ions with charge state of 2–4 and an abundance threshold of more than 150 counts, and exclusion time was set at 12 s. IDA rolling collision energy script was used for automatically adapting the CE. Mass calibration of the analyzer was achieved using peptides from digested BSA. The complete system was fully controlled by Analyst TF 1.6 (AB Sciex).

**Protein Identification**

Mass data collected during nanoLC-MS/MS were searched using a local Mascot server (Matrix Science, London, U.K.) against an in-house-generated protein database composed of protein sequences of hGR, PfGR and BSA using an in-house database generation toolbox (https://msda.unistra.fr). Searches were performed with selected modification (on each 20 encoded proteinogenic amino acids either +375.07 Da (9 or 9-BX), +374.07 Da (9-1 Da), +373.06 Da (9-BX) +357.06 Da (9 – H2O), +356.06 Da (9-BX – OH), +358.07 Da (9 – OH), +345.10 Da (9-NH2), +343.08 Da (9-BX-NH2) +327.09 Da (9-NH2 – H2O), trypsin was selected as the enzyme, carbamidomethylation of cysteine (+57 Da) and oxidation of methionine (+16 Da) were set as variable modifications, three misscleavages were tolerated and mass tolerances on precursor, and fragment ions of 20 ppm and 0.07 Da were used, respectively. Modified peptides were manually validated. Selected peptides binding sites were visualized, and distances were calculated on hGR (PDB ID: 3GRS; 2GHS) and PfGR (PDB ID: 1ONF) structure models using Chimera software.

**HPLC-MS Analysis**

LC/MS analyses were performed using an Agilent 1100 series LC coupled to a MicrOTOF-Q (Bruker Daltonics, Bremen, Germany) or to a maXis Q TOF mass spectrometer (Bruker). The mass spectrometer was operated in positive mode with a capillary voltage of 4500 V. Acquisitions were performed on the mass range of 200–1850 m/z. Calibration was performed using the singly charged ions produced by a solution of Tune mix (G1969–85000, Agilent, U.S.A.). Data analysis was performed by using Compass DataAnalysis 4.3 (Bruker Daltonics). A cross-linking reaction mixture containing GSH and PDO (or probe 9) was directly analyzed onto a HPLC connected to MicrOTOF-Q. Compounds were separated on a XBridge Peptide BEH C18 column (300 Å, 3.5 μm, 2.1 mm × 250 mm) column. The gradient was generated at a flow rate of 250 μL/min using 0.1% trifluoroacetic acid (TFA) in water for mobile phase A and ACN containing 0.08% TFA for mobile phase B at 60 °C. Phase B was increased from 5 to 85% in 45 min.

**MS Fragmentation of Collected Fractions**

Adducts synthesized in photoreactions and selected for fragmentation were purified by HPLC into specific fractions. Fragmentation of compounds and standards was performed on a hybrid electrospray quadrupole time-of-flight mass spectrometer MS (Synapt G2 HDMS, Waters, Manchester, U.K.) coupled to an automated chip-based nanoelectrospray device (Trivessa Nanomate, Advion Biosciences, Ithaca, U.S.A.) operating in the positive ion mode. The MS analysis was performed on the Synapt G2 HDMS instrument with external calibration using the singly charged ions produced by an ES-TOF tuning mix (G1969–85000, Agilent, U.S.A.). The nanoelectrospray device (Trivessa Nanomate) was set at 1.5 kV on capillary, and the pressure of the nebulizer gas was 0.55 psi. Selected ions were fragmented with a collision energy ranging from 5 to 40 eV until sufficient fragmentation was achieved.

**Collision-Induced Dissociation-Electrospray Mass Spectrometry Measurements**

Electrospray mass spectra of heme complexes were obtained with a Bruker Daltonics MicroTOF spectrometer (Bruker Daltonik GmhH, Bremen, Germany) equipped with an orthogonal electrospray (ES) interface. Calibration was performed using Tuning mix (Agilent Technologies). CID experiments were performed with a capillary exit (cone voltage) ranging from 120 to 400 V with 20 V increments. Stock solution of hematin ([FeIIIgpPIX (OH)3]3− or [FeIIIpPIX (OH)]2−) was freshly prepared from hemin (ferrirprotoclinic acid, S)-Fe(II)PPIX. The reaction was UV-irradiated for 10 min and the mixture was analyzed by HPLC-MS.
porphyrin chloride, [Fe^{II}PPIX (Cl)]^{2+} just before use in 50% ammonia. Stock solution of benzoanthone BX 4 (5 mM) was prepared in ACN, while chloroquine (CQ, 2.91 mM) and amodiaquine (AQ, 2.28 mM) were prepared in water. Hematin and the substrate (4 or CQ or AQ) were mixed together in CH_{2}CN/H_{2}O (50:50 v:v) in order to obtain equimolar concentrations of 100 μM. Prior to analyses, the samples were further diluted at 50 μM in ACN/H_{2}O/HCOOH (50:50:1 v:v:v). The sample solutions were then introduced into the spectrometer source with a syringe pump (Harvard type 55 1111: Harvard Apparatus Inc., South Natick, MA, U.S.A.) with a flow rate of 5 μl·min^{-1}. Stability responses of the heme-drug adducts obtained by ESI-CID experiments were drawn.

**Antimalarial Activity**

Antimalarial activity is represented by IC_{50}, the concentration of biomolecule at which half of the parasites are killed. The antimalarial activity of PD-ABPP was evaluated on asexual parasite stages of the chloroquine-resistant Dd2 strain of *Plasmodium falciparum*. Highly synchronised young rings (0-3h old) were treated for 72 h with various concentrations of PD-ABPP. For this, mature schizonts of highly synchronous parasite cultures were separated using 60% percoll gradient and the mature segmented schizonts were incubated for 3 hours for reinvasion. Remaining mature stages were then killed with 5% sorbitol and parasitemia adjusted to 0.5% in 1.5% haematocrit.

Parasite replication was assessed by fluorescent SYBR green staining of parasitic DNA as previously described. After 72 hours, the parasites were frozen at −80 °C overnight, followed by thawing and the DNA stained by SYBR green and the fluorescence measured on a plate reader (Promega) at 591 nm wavelength. For each well, the signal before adding SYBR green was subtracted to the signal with SYBR green to eliminate compound fluorescence. The percentage of surviving parasites compared to controls was determined as follows: \( \frac{\text{Fluo}_{X} - \text{Fluo}_{A,B,C}}{\text{Fluo}_{X} - \text{Fluo}_{D,M,S,O}} \times 100 \) where Fluo_{X} and Fluo_{D,M,S,O} are the mean fluorescence signals of parasites incubated with compound X or DMSO as a control, and Fluo_{A,B,C} the mean fluorescence signal of infected parasites exposed to high drug concentrations that kills them all to account for DNA of the starting parasites in each well (0.5% parasitemia). IC_{50} values were calculated using Prism (GraphPad, log(inhibitor) vs normalized response − Variable slope). Control parasitemia at T0 and T72 was determined by microscopic analysis of Giemsa stained blood smears to verify parasite stages and multiplication factor.

**ASSOCIATED CONTENT**

 Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacsau.1c00025.

Detailed experimental procedures, spectroscopic data, NMR and mass spectrometry analyses of the new compounds 7–11 and of various partners-probes adducts (PDF)

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**Notes**

The authors declare no competing financial interest.

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