Current and emerging target identification methods for novel antimalarials

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ABSTRACT

New antimalarial compounds with novel mechanisms of action are urgently needed to combat the recent rise in antimalarial drug resistance. Phenotypic high-throughput screens have proven to be a successful method for identifying new compounds, however, do not provide mechanistic information about the molecular target(s) responsible for antimalarial action. Current and emerging target identification methods such as in vitro resistance generation, metabolomics screening, chemoproteomic approaches and biophysical assays measuring protein stability across the whole proteome have successfully identified novel drug targets. This review provides an overview of these techniques, comparing their strengths and weaknesses and how they can be utilised for antimalarial target identification.

1. The need for new antimalarial compounds with novel mechanisms of action

Malaria remains one of the most prevalent infectious diseases throughout the world, with more than 40% of the world’s population living in malaria endemic regions (World Health Organization, 2021). In 2020, there were over 200 million cases and over 600 thousand deaths caused by malaria, predominantly due to two Plasmodium species, P. falciparum and P. vivax (World Health Organization, 2021). Of the two, P. falciparum is responsible for over 90% of human infections and represents a significant global health burden. These cases disproportionately affect developing countries, with nineteen African countries along with India carrying 85% of the malaria burden. Furthermore, the majority of deaths occur in children under 5 years of age. Efforts in malaria prevention have had mixed success. The use of insecticide treated bed nets has been effective in reducing parasite prevalence and child mortality, and has been a core aspect of malaria control programs (Prype et al., 2018). However, the development of a vaccine has had limited success, with the recently approved RTS,S vaccine (Mosquirix™) demonstrating limited efficacy, reducing clinical malaria episodes by 26–38% in children (Morrison, 2015). In the absence of a highly efficacious vaccine, management of malaria currently relies on small molecule antimalarial drugs that can quickly and effectively treat active malaria infection, prevent new infections or remove quiescent liver-stage parasites.

Currently, artemisinin-based combination therapies (ACTs) are the front-line treatment for P. falciparum infections. These combine fast acting artemisinin derivatives with complementary antimalarials that act via a different mechanism of action (Heller and Roepe, 2019). Combination therapies provided a solution to the development of resistance to previously-used antimalarials such as chloroquine, sulfadoxine and pyrimethamine throughout Africa (D’Alessandro and Buttens, 2001). However, the development of artemisinin resistance throughout South East Asia over the previous decade threatens the effectiveness of ACTs (Dondorp et al., 2009). Therefore, the development of new antimalarials with novel mechanisms of action is necessary to further reduce malaria-associated mortality and proceed towards complete eradication (Anthony et al., 2012; Phillips et al., 2017).

Many new compounds with promising antimalarial potential have recently been discovered using phenotypic high throughput screens. Phenotypic screens test libraries of compounds against whole cells in order to determine those compounds with antiparasitic activity, via a phenotypic readout such as parasite survival (Hovlid and Winzeler, 2016). These screens have been a valuable tool for the identification of new classes of antimalarials as they do not require prior knowledge of a validated biological target (Hovlid and Winzeler, 2016; Katsuno et al.,

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2. Drug target identification methods

2.1. In vitro evolution and whole genome sequencing (IVIEWGA)

A commonly used method for the identification of unknown targets involves in vitro selection of resistance to a novel compound coupled with whole genome sequencing (IVIEWGA) (Cowell and Winzeler, 2019a; Xie et al., 2018). In this method, parasites are exposed to low concentrations of antiparasitic compounds until resistant parasites are selected. The resistant clones can then be sequenced via whole genome sequencing to identify genetic mutations or copy number amplifications that confer resistance (Cowell et al., 2018; Xie et al., 2018). These mutations typically confer resistance by reducing the ability of the drug to bind to its target or by developing a compensatory mechanism through which the parasite can overcome the drug pressure (Cowell et al., 2018).

Successful IVIEWGA approaches, experimental design, and the role that it has played in antimalarial drug discovery have previously been reviewed in detail (Cowell and Winzeler, 2019b; Flannery et al., 2013; Luth et al., 2018; Nzila and Mwai, 2010; Okombo et al., 2021).

IVIEWGA has been used to explore the MoA of many novel antimalarial chemotypes. This is perhaps most evident in the mapping of the *P. falciparum* druggable genome (Cowell and Winzeler, 2018). Cowell et al. selected for in vitro resistance against 37 chemically distinct small molecules, identifying 35 genes thought to be either a drug-resistance determinant or an actual target. A number of these were new drug targets, including farnesyltransferase, dipeptidyl-aminopeptidase I and the aminophospholipid transportin P-type ATPase. IVIEWGA has also successfully identified a number of new targets for compounds now in clinical development, such as DDD107498 and DSM265, which target translation elongation factor 2 (eEF2), and dihydroorotate dehydrogenase (PfDHODH), respectively, highlighting the power of IVIEWGA as an untargeted approach for target identification (Baragana et al., 2015; Cowell and Winzeler, 2019a; White et al., 2019).

A primary advantage of IVIEWGA is that identified mutations have demonstrable phenotypic implications for parasite survival, in contrast with other target identification methods that identify direct target-protein interactions. Furthermore, identified resistance mechanisms could provide insight into the potential for resistance to develop in the field if these compounds are used clinically. However, IVIEWGA also has a number of limitations. The identification of mutations via whole genome sequencing must be validated, either by reintroduction of the mutation into wild type parasites, or an extensive mechanistic investigation to confirm the mutation identified is responsible for the resistant phenotype observed.

In vitro resistance generation can induce non-specific, multi drug resistance mechanisms that provide limited information about the specific target of the test compound. For example, four months of continuous culturing of *P. falciparum* in the presence of sub-lethal concentrations of the imidazolopiperazine KAF156 (ganaplacid), resulted in the selection of six resistant clones (Kuhnen et al., 2014; M. P. Challis et al.).

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Table 1

| Technique                          | Strengths                                                                 | Weaknesses                                                                 | Examples*  |
|------------------------------------|---------------------------------------------------------------------------|---------------------------------------------------------------------------|------------|
| **In vitro resistance selection and whole genome sequencing** | • Established technique                                                  | • Time consuming. Can take multiple months to select for resistance       | • (Baragana et al., 2015; Dorjsuren et al., 2021; Favuzza et al., 2020; Flannery et al., 2015; Krishnan et al., 2020; Lawong et al., 2021; Reader et al., 2021; White et al., 2019) |
|                                    | • Identified mutations have phenotypic implications for parasite survival and therefore are directly related to compound activity | • Indirect and non-specific resistance mechanisms can confound target identification |            |
|                                    | • Resistance-causing mutations in target proteins can define the binding site of active compounds - valuable for structure-enabled drug discovery | • Resistance-causing mutations cannot be identified for a significant number of ‘irresistible’ compounds |            |
| **Untargeted multi-omics approaches** | • Can be performed rapidly and at a medium throughput                     | • Limited to blood stage parasites                                        | • (Allman et al., 2016; Clements et al., 2020; Creek and Barrett, 2014; Giannangelo et al., 2020; Vanaerschot et al., 2020) |
|                                    | • Hypothesis generating. A metabolic profile can guide future investigation | • Rarely identifies a single target                                       |            |
|                                    | • Can allow for quick characterisation of known modes of action by comparing to antimalarials with established metabolic signatures | • Identified metabolic changes can be indirectly related to the compound target |            |
| **Affinity-based probe chemoproteomics** | • Identifies direct compound-protein interactions                          | • Potency and speed of onset of the compound can influence the ‘omic’ profile | • (Paquet et al., 2017; Penarete-Vargas et al., 2014; Siddiqui et al., 2022; Vanaerschot et al., 2020) |
| **CETSA proteomics**               | • Biophysical assay of compound-protein binding                           | • Requires the synthesis of specialised tool compounds                    | • (Dziak et al., 2019; Lu et al., 2020) |
|                                    | • Biophysical assay of compound-protein binding                           | • Differentiating between direct target binding and non-specific binding can be challenging without sufficient controls |            |
| **Limited proteolysis mass spectrometry** | • Can distinguish between allosteric and catalytic binding sites          | • Susceptible to false negatives                                       | • (Piazza et al., 2018, 2020) (Non-Plasmodium) |
|                                    | • Detect structural changes as a result of compound binding               | • Poor membrane protein coverage                                               |            |
LaMonte et al., 2016; Meister et al., 2011). Upon sequencing, mutations in one specific gene, *P. falciparum* cyclic amine resistance locus (*PfMDR1*), were found to confer resistance (Kuhnen et al., 2014). The function of *PfMDR1* is unknown, although homology models have suggested a potential role in protein trafficking (Jonikas et al., 2009). Interestingly, mutations in *PfMDR1* also conferred resistance to structurally unrelated compounds (LaMonte et al., 2016; Magistrado et al., 2016) and the resistance-causing mutations were not present in an obvious catalytic site, suggesting that *PfMDR1* is likely a general resistance determinant rather than a molecular target. A recent study has localised the imidazolopiperazine GNFI79 to the endoplasmic reticulum and demonstrated inhibition of protein transport in *P. falciparum* as well as the expansion of the ER following imidazolopiperazine treatment (LaMonte et al., 2020). However, the molecular target of the imidazolopiperazines has not yet been identified and further studies using chemoproteomic or biophysical methods may be required to identify a direct target.

Other examples of non-specific resistance mechanisms include an increased copy number of *Pf* multi-drug resistance 1 transporter (*PfMDR1*). *PfMDR1* confers multi-drug resistance via enhanced drug efflux, and therefore would provide little insight into a potential target (Blasco et al., 2017; Cowell and Winzeler, 2016). *PfATP4* mutations are another resistance mechanism which have been shown to confer resistance to multiple, chemically distinct classes of compounds including the spiroindolones, pyrazoles and the dihydroisoquinolones (Flannery et al., 2015; Spillman and Kirk, 2015). While *PfATP4* is an important drug target for the clinical candidate cipargamin (Rottmann et al., 2010), its propensity for conferring resistance suggests that it could have a compensatory function and therefore provides limited insight into the binding target of cipargamin.

Whilst IVIEWGA has proven very effective at identifying the mechanisms of action and/or resistance of promising drug candidates, it should also be noted that this approach preferentially identifies targets that are inherently prone to developing resistance, which could be detrimental to future clinical usage. For example, resistance to atovaquone can be rapidly selected for *in vitro* (Masset et al., 2007; Phillips et al., 2015), and resistance to atovaquone rapidly emerged in the field shortly after it was first introduced as a monotherapy (Looaeresuwan et al., 1996). Therefore, new strategies to expedite the early prediction of parasite resistance to new antimalarial compounds are of significant interest to the antimalarial drug discovery community. One such strategy is the minimum inoculation of resistance (MIR) parameter, which describes the potential for selection of resistance to a given candidate compound at a given concentration (Duffey et al., 2021). Obtaining an MIR involves exposing a range of parasite inocula (10^5 to 10^7 parasites) to constant drug pressure to determine the minimum number of parasites required for selection of a resistant population. The MIR parameter allows for the identification of compounds for which resistance can easily be selected. These compounds can then be deprioritised in preference for compounds which may be less susceptible to the emergence of resistance in the field. The MIR parameter is impacted by both the concentration of compound used and the parasite strain tested. Therefore, the use of the multi-drug resistant *P. falciparum* Dd2-B2 line is recommended and it is important to generate a MIR at multiple concentrations (Duffey et al., 2021). The MIR and speed of resistance selection can provide insight into the MoA of new compounds. For example, compounds for which resistance is difficult to select could possess multiple parasite targets, or have a fast onset of action.

Furthermore, ‘irresistible’ compound classes, where the specific MoA of the compound prevents the selection of resistant parasites via *in vitro* evolution methods, are becoming of increasing interest to the antimalarial drug discovery community (Yang et al., 2021). These compounds differ from ‘targetless’ compound classes, where resistant parasites can be obtained *in vitro* however, a target cannot be determined from IVIEWGA alone. Irresistible compounds are less likely to result in drug resistance when used clinically and often have a fast onset of action (Cowell and Winzeler, 2018). Therefore, the Medicines for Malaria Venture (MMV) have highlighted the development of ‘resistance-proof’ chemical scaffolds as a priority for future antimalarials (Burrows et al., 2017). Other target identification methods may be necessary for these compound classes. However, the use of mutagenic agents such as ethyl methanesulfonate to increase the likelihood of selecting resistant parasites has seen success in generating resistance-causing mutations for compounds where traditional IVIEWGA experimental designs have failed (Gisselberg et al., 2018). Furthermore, hypermutator parasite lines that possess mutations in the DNA polymerase delta, which ablates their proofreading activity, have been shown to increase the rate of DNA mutation (Honma et al., 2014). These lines have been successfully used to identify putative targets for the antimalarial Salinipostin A, after traditional IVIEWGA approaches failed to identify resistance-causing mutations (Yoo et al., 2020). While hyper-mutating strategies can increase the odds of identifying resistance-causing mutations, extensive validation of identified mutations is necessary to ensure they are responsible for the resistance phenotype, and are directly related to the compound’s mechanism of action.

### 2.2. Systems biology multi-omics approaches

Untargeted omics techniques, such as metabolomics, proteomics and transcriptomics, can provide an untargeted overview of cellular systems by the simultaneous detection and quantification of small molecules, proteins and transcript levels in a biological system (Dunn et al., 2012; Scalbert et al., 2009). By comparing drug-treated samples to untreated controls, perturbations in cellular processes resulting from drug treatment can be identified. This can provide insight into the MoA of novel compounds and in some limited instances, identify specific targets.

Of the different omics techniques, untargeted metabolomics is best positioned for identifying direct effects of drugs on parasite metabolism, whereas transcriptomics and proteomics are more likely to detect downstream or secondary responses. Because of this, untargeted metabolomics approaches have been successfully applied to confirm identified antimalarial targets, with studies into the MoA of atovaquone providing a good example of how untargeted metabolomics can demonstrate a specific MoA (Allman et al., 2016; Dickerman et al., 2016). Atovaquone is a direct inhibitor of the bc1 complex, an essential component of the mitochondrial electron transport chain (Fry and Pudney, 1992). Inhibition of the bc1 complex impairs *de novo* pyrimidine synthesis through the indirect inhibition of *PfDHODH* and this produces a clear metabolic signature, with a rapid and robust accumulation of the precursors for pyrimidine synthesis, N-carbamoyl-L-aspartate and dihydroorotate, detected following atovaquone treatment (Allman et al., 2016). This clear metabolic signature is consistently seen with other inhibitors of pyrimidine synthesis, such as DSM265, and has been used to identify *PfDHODH* as a target of MMV020439 and MMV007571 (Coteron et al., 2011; Creek et al., 2016; Dickerman et al., 2016).

Metabolomics can also be used to characterise the MoA of novel compounds with higher throughput than other target identification methods, by comparing the metabolic profile of test compounds with those of known compounds. Two studies into the MMV Malaria Box were able to predict a substantial number of target pathways for compounds with previously unknown MoA (Allman et al., 2016; Creek et al., 2016). One study simultaneously investigated the mode of action of 90 antimalarial compounds, with over half of the tested compounds demonstrating significant metabolic perturbations. Of these compounds, nearly one quarter caused an increase in N-carbamoyl-L-aspartate and dihydroorotate, indicating that they inhibit pyrimidine synthesis, while 13 compounds possessed a metabolic signature similar to artemisinin and 6 shared a metabolic signature with chloroquine. This allows for the quick characterisation of novel compounds, providing rapid insight into their potential MoA. However, it is important to consider that while similar metabolic profiles could indicate similar MoA, they do not necessarily have the same biological target. For instance, it was
hypothesised that those compounds that clustered with artemisinin may share its potent, fast acting profile, and therefore the observed metabolic perturbation was likely due to rapidly mediated cell death, rather than an indicator of a specific drug target or pathway (Creek et al., 2016).

A limitation of typical metabolomics workflows is the fact that they only detect small molecules and therefore, are not useful for compounds which do not interfere with parasite metabolism. Metabolomics has been combined with other ‘omics’ approaches, such as transcriptomics or proteomics to overcome this limitation (Birrell et al., 2020; Gianangelo et al., 2020; van Brummelen et al., 2009). These complimentary approaches allow for a greater range of biomolecules to be detected, providing more scope for the identification of changes that can provide insight into the MoA. For instance, a recent investigation into synthetic oxonide antimalarial combinations performed at later time points identified secondary perturbations in pyrimidine biosynthesis, the Kennedy pathway, translation regulation and the ubiquitin-proteasome system, providing a comprehensive analysis of both primary and secondary mechanisms of oxonide-induced parasite death.

Another limitation of typical metabolomics-based approaches is that while they often provide valuable mechanistic information, they often identify a target pathway or signature rather than the specific drug target, and they cannot measure direct drug-target binding. Systems biology-based approaches to investigate compound mechanisms can also be confounded by many experimental design factors, including compound concentration, duration of incubation and parasite lifecycle stage. Designing experiments with multiple distinct time points and concentrations can assist with identifying drug-specific perturbations. Nevertheless, untargeted metabolomics, proteomics and transcriptomics approaches provide useful and unbiased characterisations of the impact of novel drug candidates on parasite biochemistry, enabling the generation or confirmation of hypotheses regarding their MoA.

2.3. Chemoproteomic approaches to target identification

Small molecule affinity-based probes have been widely used for the identification of novel drug targets and new biological pathways (Heal et al., 2011; Su et al., 2013). Affinity-based probes require functionalisation of the compound of interest with chemical handles that allow the compound to be enriched, along with its bound protein target(s) (Kawatani and Osada, 2014). Recently, novel antimalarial compounds with unknown MoAs have been functionalised to allow for the identification of new drug targets (Ismail et al., 2016; Paquet et al., 2017). The study into MMV390048 is an example where an affinity-based probe was used to successfully identify P. falciparum phosphatidylinositol 4-kinase (PfPI4K) as a new target for a novel antimalarial compound (Paquet et al., 2017). An active analogue, MMV666845, was covalently immobilised on Sepharose beads and used as bait to capture protein targets from a cell lysate. To distinguish between specific and non-specific binding, this pull-down approach was also performed in the presence of the active parent compound MMV390048. The ‘free’ MMV390048 outcompetes the probe MMV666845 for the protein target, resulting in less enrichment of the target protein compared to non-specific binding proteins. This allows the target to be distinguished from those proteins with non-specific binding. This approach identified PfPI4K as the only specifically-bound protein and was supported by in vitro generation of MMV390048-resistant parasites, which identified mutations in PfPI4K.

A convenient form of affinity-based probes are those based on copper catalysed ‘click chemistry’ reactions (Speers and Cravatt, 2004). ‘Clickable’ azide and alkyne functional groups can react, forming stable triazole conjugates (Fig. 1. A). These ‘clickable’ functional groups can be synthetically introduced onto the compound of interest and, after exposing these functionalised probe compounds to either live cells or a cell lysate, they can be reacted with the reciprocal functional group which is typically immobilised on beads. This reaction forms a covalently-linked triazole conjugate, allowing for the probe to be pulled out of the complex proteome, along with any interacting protein(s) that can subsequently be detected via mass spectrometry (Fig. 1. B). ‘Clickable’ probes have been used to identify protein targets for endoperoxide antimalarials including artemisinin and synthetic oxonide compounds (Ismail et al., 2016; Jourdan et al., 2019; Siddiqui et al., 2022; Wang et al., 2015). These studies identified proteins from a range of pathways including haemoglobin digestion, antioxidant defence systems, glycolysis, nucleic acid and protein biosynthesis, demonstrating the expected promiscuous nature of artemisinin binding within the parasite as a result of the radical chemistry involved with the endoperoxide bond cleavage.

A limitation of affinity-based probes such as MMV666845, is that the interactions with their target proteins are non-covalent and therefore can be transient. One approach to overcome this challenge is the use of bifunctional probes that can covalently bind to target proteins through photoactivatable functional groups such as phenyl azides or diazirines, as well as possessing a chemoreactive group suitable for enrichment. A novel, non-covalently binding antimalarial, albitalozium, was investigatd with photoactivatable bifunctional probes which could covalently bond to target proteins to aid in the determination of specific drug-protein interactions (Penarete-Vargas et al., 2014). These bifunctional probes showed good specificity, identifying eleven proteins as interactors and providing a shortlist of promising targets for further validation. This demonstrated the scope for development of advanced probes that improve the reliability and versatility of this approach for compounds that don’t act through covalent binding.

These studies demonstrated how chemoproteomic approaches can identify binding targets for antimalarial compounds with unknown MoAs, in an untargeted fashion, however, a number of factors limit their use. Firstly, this approach requires the time-consuming synthesis of specifically designed chemical probes with the desired functionalities. This is further complicated by the need for good knowledge of the structure activity relationships of the compound of interest, to ensure that the probe modifications don’t interfere with compound activity or target specificity. Therefore, it is not suitable for high throughput screening applications and likely will only be applicable for compounds showing significant promise. Secondly, approaches that are dependent on affinity to identify target proteins can be susceptible to identifying non-specific interactions and co-purified proteins (Gingras et al., 2007). While this can potentially confound the results, sufficient negative controls, particularly with competitive binders, can ameliorate this. Furthermore, the identification of co-purified proteins can provide context for the biological role of the identified target. This can be particularly important in the context of P. falciparum, where a significant proportion of the genome has no functional annotation.

Databases of commonly identified contaminants, such as the Contaminant Repository for Affinity Purification Mass Spectrometry Data (CRAPOme) can help to identify non-specific protein binding for Homo sapiens and Saccharomyces cerevisiae data (Melicheruu et al., 2013). However, a similar P. falciparum database is not currently available, placing more importance on having sufficient negative controls and replicates to help discern non-specific interactions from real targets. This is highlighted by the requirement of a chemoproteomic approach to directly compare the alkylation profiles of endoperoxide antimalarial probes, where the use of advanced mass spectrometry techniques and extensive negative controls allowed for the high confidence identification of new alkylation targets for endoperoxide antimalarials (Siddiqui et al., 2022).

Overall, these studies demonstrate the power of chemoproteomics as an untargeted technique for target identification, and highlights the importance of direct approaches that can distinguish between molecular
targets and indirect resistance mechanisms, provided strategies to distinguish between target binding and non-specific binding are employed effectively.

### 2.4. Protein stability target identification

When proteins are heated they denature, unfold and form insoluble aggregates (Varela et al., 2019). However, the preferential binding of a ligand to a target protein can make that protein more resistant to denaturation, resulting in a higher proportion of ligand-bound protein remaining in its soluble form following a thermal challenge. After a thermal challenge, soluble proteins can be separated from denatured aggregates by centrifugation and ligand-stabilised proteins can be identified by an apparent increase in abundance compared to a negative control. This concept has been applied to a number of targeted thermal shift style assays utilising Western blot or fluorescent detection methods (Martinez Molina et al., 2013; Martinez et al., 2018). However, these methods typically require either purified protein, or antibodies directed at a suspected target, and therefore are more suitable for the validation of suspected targets rather than the identification of unknown MoAs (Martinez et al., 2018; McNulty et al., 2018; Shaw et al., 2018). The recent incorporation of quantitative proteomics techniques (Franken et al., 2015; Martinez Molina et al., 2013) has allowed these thermal shift assays to be simultaneously performed across whole proteomes, therefore making it possible to identify previously unknown protein targets from a complex proteome. These whole proteome approaches are typically referred to as Cellular Thermal Shift Assays (CETSA).

CETSA has two main variants: the thermal proteome profiling (TPP) or ‘melt curve’ approach, whereby the temperature of the heat challenge is varied in the presence of a consistent concentration of compound (Franken et al., 2015), and the isothermal dose response (ITDR), which employs increasing compound concentrations with a single temperature (Fig. 2) (Dziekan et al., 2019; Franken et al., 2015). In TPP CETSA the thermal melting temperature ($T_{m}$) increases in proteins bound by the ligand, resulting in an increase in the amount of soluble protein remaining following heating. In ITDR CETSA, as the concentration of ligand is increased, more protein is protected from the thermal challenge, resulting in an increase in soluble protein proportional to the increase in ligand.

Recently, a TPP CETSA proteomics study in *P. falciparum* was performed in order to identify binding targets for quinine (QN) and mefloquine (MQ), two antimalarials with previously unknown MoAs. QN was found to selectively stabilise the purine nucleoside phosphorylase (PfPNP) in a dose-dependent manner in both lysate and intact cells.

![Fig. 1](image-url). A. Copper catalysed click chemistry reaction between azide and alkyne moieties. B. Theoretical chemoproteomic pull down. A novel compound modified with an alkyne moiety interacts with an unknown target in a complex proteome. The alkyne interacts with an immobilised azide, forming a covalent triazole conjugate which allows the target protein to be isolated and subsequently identified via mass spectrometry.

**Fig. 1.** A. Copper catalysed click chemistry reaction between azide and alkyne moieties. B. Theoretical chemoproteomic pull down. A novel compound modified with an alkyne moiety interacts with an unknown target in a complex proteome. The alkyne interacts with an immobilised azide, forming a covalent triazole conjugate which allows the target protein to be isolated and subsequently identified via mass spectrometry.

![Fig. 2](image-url). A. Example melting curve for thermal protein profiling (TPP) CETSA. $\Delta T_{m}$ represents the increased thermal melting temperature of the target protein in the presence of the ligand. B. Example stabilisation curve for an isothermal dose response (ITDR) CETSA. Red arrow represents the increased stability of the target protein in the presence of increasing drug concentrations following a heat challenge. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)
Stabilisation of PfPnP was also observed in lysate samples following MQ treatment, albeit to a lesser extent (Dziekan et al., 2019). This assay was supported by other biochemical assays and co-crystal structures in order to confirm the binding of quinine to PfPnP. It was also demonstrated that MQ can bind to PfPnP, however, the therapeutic relevance could be limited due to lower potency.

TPP approaches to target identification have also been applied to other apicomplexan parasites, including Toxoplasma gondii. The calcium dependent protein kinase 1 (CDPK1) was identified as a target of a novel antiparasitic compound termed ENH1 although, follow up validation studies did identify ENH1 polypharmacology, making it difficult to discern the primary mechanism of ENH1 activity (Herneisen et al., 2020). ITDR style CETSA approaches have also been recently used to identify previously unknown targets of the pan-kinase inhibitor staurosporine TRiC/CCT as a target of the antihistamine clemastine (Lu et al., 2020). By focusing on increasing sample replicates, rather than having multiple temperatures, Ball et al.’s isothermal shift assay (ITSA) identified 51 new kinase targets for staurosporine, a significant improvement on previously performed TTP CETSA studies (Savitski et al., 2014). The ITSA method also reduced the total number of samples required for each compound tested, ultimately reducing analysis time and increasing throughput, which could make mass spectrometry-based thermal shift approaches more amenable to routine application.

Untargeted CETSA proteomics can identify potential protein binding targets from cell lysates and whole cell samples, making it a powerful technique for identifying targets of compounds with unknown MoAs. CETSA is also advantaged by the fact that it does not rely on the time-consuming synthesis of specific probes or generation of resistant parasites and therefore can be performed relatively rapidly (Dziekan et al., 2020). However, it is important to consider the possibility that off-target binding can occur at high concentrations. Therefore, subsequent validation with targeted methods is necessary to confirm the binding and the biological relevance of identified targets. Current data analysis techniques for CETSA proteomics have also been thought to result in high levels of false negative identifications, largely due to the fact that they typically rely on a significant change in a single parameter such as melting point or protein abundance (Childs et al., 2019). New data analysis techniques such as the nonparametric analysis of response curves (NPARC) could help improve target identification for TPP CETSA (Childs et al., 2019). NPARC takes advantage of functional data analysis and nonlinear regression, rather than relying on changes in a single parameter, to identify stabilised proteins with non-canonical melting profiles, reducing the number of false-negative identifications and increasing the theoretical coverage of TPP CETSA. Current CETSA proteomics approaches also have difficulty identifying membrane proteins due to the detergent-free cell extraction procedures used (Franken et al., 2015; Rawlings, 2016). While there are some studies exploring the use of detergents in CETSA protocols (Franken et al., 2015), detergents are expected to alter protein structure and stability, and more development is required before membrane proteins can be reliably identified. Finally, there are examples of compounds that do not induce any changes in thermal stability in their validated targets, highlighting how CETSA proteomics may not be appropriate for all compounds (Becher et al., 2016; Savitski et al., 2014). Despite this, CETSA proteomics can be a powerful tool for the target identification of novel compounds with unknown MoAs, particularly when combined with targeted validation assays.

Denaturation methods other than heating have also been used to identify drug targets by measuring changes in protein stability. Stability of Proteins from Rates of Oxidation (SPROX) is a method that induces protein denaturation with hydrogen peroxide in order to test protein stability and its subsequent stabilisation by ligands in a similar way to TPP-CETSA (Strickland et al., 2013). SPROX has recently been applied to antimalarial drug target discovery. Performed in concert with TPP-CETSA, SPROX identified the essential Plasmodium protein chaperonin TrIC/CCT as a target of the antihistamine clemastine (Lu et al., 2020).

Other methods, such as limited proteolysis coupled mass spectrometry (LiP-MS) and Drug Affinity Responsive Target Stability (DARTS) use limited proteolytic digestion to identify proteins stabilised by the ligand in question. These methods rely on applying proteases with broad specificity, such as proteinase K, to either whole proteomes or purified proteins for a short period of time (Lomenick et al., 2009; Schopper et al., 2017). The limited exposure of the proteases ensures that proteolysis is directed by protein structure, with exposed proteolysis sites digested first. When a ligand binds to a target site, it prevents proteolysis of the target site and results in a differential peptide pattern which can be detected via MS. This approach has previously been used to demonstrate subtle changes in secondary protein structures, structural changes induced by allostery and identify new small molecule-protein interactions (Feng et al., 2014; Geiger et al., 2016). LiP-MS has been previously validated with the antifungal drug cerulenin, successfully identifying its known target, Fatty Acid Synthase 2 (FAS2) from over 2500 identified proteins (Pliazza et al., 2016). LiP-MS has historically been limited by the difficulty of detecting LiP-MS sites against a complex peptide background (Soh et al., 2007). However, LiP-MS and DARTS have the potential to provide another proteome-wide target identification method for novel antimalarial compounds with the ability to provide unique insight that is complementary to currently established methods.

3. Prospects and challenges

Each of the untargeted techniques detailed in this review have their own strengths and limitations that can affect how they can be applied. IVIEWGA and untargeted metabolomics approaches both benefit from identified mechanisms having clear, phenotypic implications for the parasite. However, neither method directly measures target engagement and therefore both identified resistance mechanisms and metabolic perturbations could be indirectly related to the compound’s target. In contrast, chemoproteomic and CETSA proteomic approaches can identify specific drug-protein interactions, however, require validation to confirm that identified ‘hits’ have biological relevance. The ease of application of these methods can also be a limiting factor. IVIEWGA is time consuming, often requiring multiple months of continuous parasite culture and can be particularly challenging for compounds with pleiotropic effects or a fast onset of action. Traditional affinity-based chemoproteomic approaches require the synthesis of specific probe compounds, which requires medicinal chemistry expertise and can also be a time-consuming process.

It should be noted that all of the methods discussed in this review take advantage of ‘omic’ style approaches to identify specific targets from within a whole genome or proteome. While the ability to investigate the whole genome/proteome is undoubtedly powerful, it can be limited by the time and cost associated with performing large experiments reliant on sophisticated instrumentation such as mass spectrometers. Whilst the cost-efficiency of these omics approaches has improved in recent years, this barrier can encourage the design of experiments with limited samples, reducing the number of concentrations, time points or controls tested. In turn, this can increase the likelihood of false positive target identifications from these untargeted methods. All the methods discussed in this review are susceptible to false-positive ‘hits’ being identified and this highlights the crucial need for extensive target validation following the identification of promising targets. Approaches that can directly measure compound-target binding, such as surface plasmon resonance (SPR) based assays, targeted CETSA approaches and x-ray crystallography can all provide confidence in direct compound-target binding (Maveyraud and Mourey, 2020; Molina et al., 2013; Schneider et al., 2015). Furthermore, using multiple target identification methods in parallel can also provide confidence in targets that are consistently identified. Successful target identification programs have often combined multiple approaches, with complimentary
techniques confirming both target engagement and the phenotypic relevance of identified targets (Milne et al., 2022). For example, the plasmepsin inhibitors WM4, WM5 and WM382 utilised IVIEWGA to originally identify plasmepsin X as a potential target, and demonstrated that WM382 specifically bound to both plasmepsin IX and X through targeted chemoproteomic and CETSA assays (Favuzza et al., 2020). However, one caveat is that the level of expertise required to perform multiple untargeted identification techniques for a single target identification program can be restrictive, and therefore, collaboration is critical (Cowell and Winzeler, 2018).

It should also be noted that these techniques have typically been applied to investigate compounds with a blood stage phenotype, and have seen limited use in the sexual and liver stages. The recent development of high-throughput screens for both liver stage and transmission inhibitors will provide new lead compounds which will require target identification methods that have been optimised to those stages of the lifecycle (Delves et al., 2016; Swann et al., 2016). Techniques that rely on biophysical changes, such as CETSA proteomics and LiP-MS, should be easily transferrable to other parasite stages. However, further optimisation of sample preparation procedures is likely necessary.

4. Concluding remarks

The use of target identification methods greatly assists the development of new antimalarial compounds identified in phenotypic high throughput screens. Currently-used techniques such as IVIEWGA and untargeted metabolomics profiling have been successful in providing insight into the MoA of multiple compounds currently in preclinical development, fast-tracking their progress through the antimalarial development pipeline. The emergence of affinity-based chemoproteomics, CETSA and SPROX proteomics, and the potential for approaches such as LiP-MS to be used for target identification, can improve the options available for drug discovery programs to identify new targets.

There remains a number of outstanding questions regarding the scope for further development of these techniques. They are yet to be applied in typically understudied stages of the Plasmodium lifecycle and there are still opportunities to improve their ability to delineate off target or indirect effects from biologically relevant targets directly involved in the MoA. Simplifying these methods to allow for their incorporation into high throughput screening approaches and enabling earlier access for drug discovery projects may be an important area of development. The early identification of biologically relevant targets can assist with fast compound development and the triaging of promising candidate compounds early in the drug discovery process. In the context of P. falciparum drug target identification, the development of a database of common off-target or non-specific binding proteins will assist in distinguishing between off target or indirect effects and finally, the emergence of new methods such as LiP-MS, DARTS and SPROX open up exciting new approaches to antimalarial target identification that will provide alternate avenues for identifying new biologically relevant targets. Overall, it is worth noting the importance of the combined use of multiple techniques in parallel to overcome the individual weaknesses of each method, and the subsequent validation of novel targets is crucial for the advancement of new compounds that are urgently needed to combat growing antimalarial drug resistance, and progress towards malaria elimination.

Declaration of competing interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: Darren Creek reports financial support was provided by National Health and Medical Research Council.

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Abbreviations

CETSA Cellular thermal shift assay
E.coli Escherichia coli
ITDR Isothermal dose response
IVIEWGA In vitro evolution and whole genome sequencing
LiP-MS Limited proteolysis mass spectrometry
MoA Mechanism of Action
MS Mass spectrometry
Pf Plasmodium falciparum
PfATP4 Plasmodium falciparum P-type ATPase 4
PfCARL Plasmodium falciparum cyclic amine resistance locus
PfDHODH Plasmodium falciparum dihydroorotate dehydrogenase
PfMDR1 Plasmodium falciparum multidrug resistance 1
PfPI4K Plasmodium falciparum phosphatidylinositol 4-kinase
PPNP Plasmodium falciparum purine nucleoside phosphorylase
TPP Thermal proteome profiling

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