Collaborative virtual screening to elaborate an imidazo[1,2-a]pyridine hit series for visceral leishmaniasis

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Abstract: An innovative pre-competitive virtual screening collaboration was engaged to validate and subsequently explore an imidazo[1,2-a]pyridine screening hit for visceral leishmaniasis. In silico probing of five proprietary pharmaceutical company libraries enabled rapid expansion of the hit chemotype, alleviating initial concerns about the core chemical structure while simultaneously improving antiparasitic activity and selectivity index relative to background cell line. Subsequent hit optimization informed by the structure activity relationship enabled by this virtual screening allowed thorough investigation of the pharmacophore, opening avenues for further improvement and optimization of the chemical series.
Introduction

Leishmaniasis is generally considered to be one of the most neglected tropical diseases and has strong links with poverty.\textsuperscript{1–3} It is a complex vector-borne disease caused by more than 20 species of the protozoan genus \textit{Leishmania} with symptoms ranging from localized skin ulcers to a lethal systemic disease. Of the four leishmaniasis syndromes, visceral leishmaniasis (VL) represents the greatest burden to global health, with data from 2015 suggesting it is responsible for almost 1.4 million DALYs (disability adjusted life-years) lost, representing almost 6\% of all DALYs caused by diseases on the World Health Organization’s list of neglected tropical diseases (NTDs).\textsuperscript{4} Caused by infection with \textit{L. donovani} and \textit{L. infantum} parasites transmitted by phlebotomine sand flies, visceral leishmaniasis manifests in patients as swelling of spleen and liver, renal dysfunction and, if left untreated, death. Current estimates suggest an annual new infection rate for VL of 50,000 to 90,000 cases, with most cases found in eastern Africa, Brazil and India.\textsuperscript{2,5–8}

Although treatments for VL exist, these medications are rarely fit for purpose, with major issues linked to resistance, geographic variability in efficacy, toxicity, availability and affordability, and often long and painful treatment periods requiring many weeks of hospitalization.\textsuperscript{9–13} As a result, the past decade has seen increased efforts to develop new treatments for \textit{Leishmania} related syndromes such as VL, with a primary focus on delivering pre-clinical candidates with improved profiles (broader geographic utility, reduced or no toxicity, oral dosing, up to 10 day treatment period).\textsuperscript{5} Target product profiles for these next generation VL treatments have been published and discussed widely.\textsuperscript{14,15}

A combination of target-based and phenotypic-based drug discovery approaches have been applied extensively to identify new chemical matter for anti-leishmanial discovery projects, as well as the kinetoplastid field in general.\textsuperscript{9,16–18} Despite some excellent progress in target based approaches, these often suffer from poor translation into efficacy in \textit{in vitro} infection models,\textsuperscript{19} demonstrating the importance not only of a relevant mechanism of action but also the correct physicochemical properties to access the parasites once intracellular infection has taken hold. The phenotypic approach bypasses this issue by engaging cell-infection assays as the primary screen, however, these approaches often suffer from a combination of a low hit rate and identification of a high percentage of unsuitable chemical matter, with many hit lists comprising compounds rich in PAINS,\textsuperscript{20} toxicophores and intractable chemistries. Nevertheless, these approaches have yielded an interesting pipeline of novel potential treatments for VL, some of which have progressed to the early stages of clinical or preclinical evaluation.\textsuperscript{21} These include CRK12 inhibitor GSK3186899,\textsuperscript{22} the \textit{Leishmania} proteasome inhibitors GSK3494245 and LXE-408,\textsuperscript{23,24} as well as DNDI-6174, DNDI-6148 and DNDI-0690, compounds identified via phenotypic drug discovery efforts.\textsuperscript{25}

Despite this promising situation, the reality of clinical-level attrition in drug discovery R&D dictates that if the global health community are to ensure delivery of new, improved treatments to VL patients, further new chemical entities
(NCEs) capable of tackling *Leishmania* infections still need to be identified and developed.\(^{26,27}\) Alongside the traditional target and phenotypic approaches mentioned above, recent reports have highlighted a number of efforts to identify novel anti-leishmanial NCEs via repurposing of existing drugs and clinical stage compounds,\(^{28}\) combined scaffold and parasite hopping,\(^{29}\) the screening of natural products,\(^{30}\) as well as investigation into host-mediated treatments for *Leishmania* infections.\(^{31}\)

We report herein on early discovery efforts around an anti-leishmanial chemotype derived from the output of a kinetoplastid high throughput screening (HTS) campaign. Since 2012 we (DND\(i\)) have tested over 2.2 million compounds in high throughput high content screening against kinetoplastid parasites including *Leishmania donovani* and *Trypanosoma cruzi*, the causative agent of Chagas disease,\(^{32}\) searching for novel and attractive small molecule starting points for drug discovery campaigns.\(^{5}\) Seeking to improve the efficiency and throughput of the triaging of results from these phenotypic screening efforts led to the creation of the NTD Drug Discovery Booster, a pre-competitive virtual screening model comprising key players from pharmaceutical R&D.\(^{33}\)

**NTD Drug Discovery Booster**

In the traditional approach to initial exploration of chemical space around a new HTS hit, commercial compound catalogues are probed for similar molecules and small sets of closely designed analogues are synthesized for testing.\(^{34}\) As an alternative to this, we engaged an innovative *in silico* approach, termed the NTD Drug Discovery Booster (“Booster”). The Booster process engages collaborative rounds of virtual ligand-based screening across multiple large, proprietary pharmaceutical company databases to identify and test both close analogues to the hit, as well as structurally differentiated compounds fitting the same base pharmacophore. The hit structure (“Seed”) is shared with scientists at each partner pharmaceutical company, *in silico* similarity searches are performed probing the proprietary collections of each partner, and up to 100 prospective “similar” molecules per partner are screened against the parasite(s) of interest. The definition of “similar” is independently determined by each partner, and could be focused on both SAR annotation as well as more general pharmacophore-led exploration, and could involve the use of both commercial and proprietary computational approaches. Sharing the results of these experimental screens with the providing partners, as well as anonymized sharing of the structure of the most improved molecule from the entire collaborative screening endeavor (“Improved Hit”), allows a second, more refined iteration of *in silico* screening to be performed, again providing approximately 100 molecules per partner. Repetition of this iterative screening / sharing process continues until improvements cease to be made, or until SAR around a chemotype emerges.

The use of both orthogonal *in silico* techniques as well as orthogonally differentiated libraries, previously demonstrated to have minimal overlap,\(^{35}\) maximizes both the richness of the SAR explored around the initial seed phenotype, and increases the chances of identifying an interesting change of chemotype with similar or improved...
anti-parasitic activity, i.e. a scaffold hop. Furthermore, these newly identified scaffold hops can be subsequently mined using the same Booster approach, thus maximizing the value of the hits coming from HTS.

Results and discussion

Hit identification

Compound 1 was identified as a hit with anti-trypanosomal activity from a high throughput screen against *Trypanosoma cruzi*. Follow up confirmation of activity validated both the anti-trypanosomal activity of 1 in a *T. cruzi* intracellular infection assay (U2OS as host cell) as well as demonstrating the compound’s efficacy against *Leishmania donovani* in an intracellular infection assay (THP1 as host cell) (Fig 1). As mentioned previously, we are wary of the high hit rate of intractable, non-drug-like and/or PAINS-like compounds in the kinetoplastid literature, and our initial analysis of the 1 chemotype flagged up the 2-pyridyl 3 amino substituted imidazo[1,2-a]pyridine core as a PAINS motif. Further investigation of the literature around 1 identified earlier reports about a related structure with moderate potency against the related leishmania parasite *L. major*, one of the causative parasitic agents of cutaneous leishmaniasis. In this report the chemotype did not contain the 2-pyridyl motif nor the amino substituent at the 3 position of the imidazo[1,2-a]pyridine, but rather a 2,3 biaryl substitution pattern. In this instance the structure is, therefore, not classified as a PAINS motif, however the reported SAR demonstrated potencies around an order of magnitude lower than those observed for 1, despite the authors using an axenic promastigote assay which circumvents the additional challenge for compounds to access intracellular parasites. In addition, we found examples of launched drugs which contain the imidazo[1,2-a]pyridine core from 1, including the so-called Z-drugs Zolpidem and Alpidem, as well as reports of structures similar to 1 confirmed as hits in other infectious disease screens such as *Mycobacterium tuberculosis* and vaccinia virus.

Combining the above findings with the nature of the assay used to identify the anti-leishmanial effect of the compound (high content imaging with parasite count) and the fact that the compound demonstrated anti-parasitic activity in more than one assay and against multiple related kinetoplastid parasites, we decided to cautiously investigate further. We were aware from the literature of the potential for a metal-chelation-driven pharmacological effect due to the bidentate heteroaryl-heteroaryl system in the core, indeed it could be hypothesized that this is the source of PAINS-like activity in other assays. Therefore, a key step for further investigation of 1 was to gain confidence via SAR and orthogonal assays that the observed anti-parasitic activity was unlikely to be driven by metal chelation, and that the PAINS flag could be legitimately discounted. The two key read-outs we targeted to achieve this were a) the ability of the chemotype to achieve a broader selectivity index for *Leishmania donovani* relative to the background host cell, and b) clear evidence that altering the core away from the PAINS-like chemotype maintained anti-parasitic activity, as suggested by the earlier paper from Marhadour et al.
Hit elaboration and validation via collaborative *in silico* screening

Compound 1 was submitted as a “seed” to five partners of the NTD Booster consortium: AstraZeneca plc, Celgene Corporation, Eisai Co., Ltd, Shionogi & Co., Ltd, and Takeda Pharmaceutical Company Ltd. Since the pre-competitive mechanism of the Booster project requires anonymity and a high degree of confidentiality on *in silico* methods and library composition between partners, identities of the partners are presented as blinded in this study as “Companies A-E”. §§

The partners performed two successive iterations of ligand-based virtual screening against their proprietary compound collections: cycle A and cycle B (see fig 1). Each distinct library was probed using a partner-specific *in silico* similarity search approach as described in Table 1 (see supporting information for detailed description). The compounds identified via these virtual screening efforts were investigated in a high content cell-based *L. donovani* infection assay using THP1 host cells. Assay read-out from this high content approach demonstrated the compound’s efficacy at total parasite clearance as well as host cell cytotoxicity, yielding an anti-parasitic IC50, a host cell CC50 and a selectivity index (SI) between the two read-outs (SI = host cell IC50 / parasite IC50; see supporting info for more details).

Compounds with a SI > 5 were considered as demonstrating legitimate anti-parasitic activity. Results of this process are shown in Fig 1. The first sequence of virtual screening (Fig 1 A) identified a total of 367 compounds from 4 partners, of which 25 demonstrated at least some degree of potency against *L. donovani* and shared the core imidazo[1,2-a]pyridine motif. Interestingly a new batch of compound 1, which displayed similar potency - cytotoxicity profile to the batch from the original HTS, appeared within this set (Fig 2 A). Compound 2 also appeared within this set, which demonstrated the potential of a new vector for exploration at the 8-position of the imidazo[1,2-a]pyridine core, as well as demonstrating efficacy with no background cell cytotoxicity. Parasitology data on each company’s selection of compounds was fed back to the respective partner organizations. Parasitology data and the structure of compound 2 was shared with all partners and a second round of virtual screening was then completed using this elaborated parasitology data (fig 1 B). The second round of *in silico* screening yielded a further 290 compounds from 5 participating organizations, of which 16 contained the imidazo[1,2-a]pyridine core and demonstrated some level of efficacy against *L. donovani*. Retrospective analysis of these two rounds of *in silico* screening demonstrated the power and versatility of this approach to hit to lead elaboration. Of the 657 compounds identified via *in silico* screening, almost 45% of the compounds represented a scaffold hop, defined in this instance as a compound not containing a 2-substituted imidazo[1,2-a]pyridine as the core. The hit rate of actives within this set of scaffold hops was only marginally lower than for those “on-scaffold” compounds which contained the original chemotype (11.5% compared to 15.4%) - yet these more structurally ambitious scaffold hop compounds would arguably not be included in a traditional hit to lead campaign due to the inherent increase in synthetic complexity of scoping multiple chemotypes. Compound 3
is a key example of such a scaffold hop originating from this approach, demonstrating significant potencies against both *T. cruzi* and *L. donovani* whilst representing a significant deviation away from the imidazo[1,2-a]pyridine core.

Combining the parasitology data for the key active compounds containing the original imidazo[1,2-a]pyridine core structure of 1, we collated data to form a hit series and move forward into detailed investigation and targeted analoguing as detailed below. An initial survey of the data generated by these rounds of virtual screening resulted in a number of initial findings: Firstly, there appeared to be at least 5 vectors around the core which tolerated changes whilst retaining anti-leishmanial activity (positions 2, 3, 6, 7 and 8). Secondly, although certain structural changes in the chemotype induced significant cytotoxicity against the host THP1 cell line, we identified key examples which demonstrate acceptable differentiation between anti-parasitic activity and cell based cytotoxicity, with the most selective compounds achieving a selectivity index (SI) of > 20 (4, 5, 6, 7). Analysis of the anti-parasitic activity and cell line cytotoxicity curves and high content images clearly demonstrated concentration points for most compounds where a full antiparasitic effect was observed with no cytotoxic observation on the background cell line (see Fig 2B, C). Thirdly, we were able to identify compounds which achieved an antiparasitic effect with no background cell cytotoxicity whilst removing the 2-pyridyl functionality at the 2 position of the core (8), removing some suspicions about a metal-chelation driven SAR. These findings convinced us that the cytotoxicity observed for the series of compounds was not uniquely responsible for the observed anti-parasitic effect, and that the series merited further investigation. Finally, we also observed that the anti-leishmanial SAR around the scaffold was potentially non-additive, with the impact of certain functionality changes being influenced by substitution patterns elsewhere in the structure (Fig 2A). This appeared particularly true for substitutions of the 6 and 7 position of the core imidazo[1,2a]pyridine, as demonstrated by compounds 4-6, where the impact of a 6-methyl group improves potency when the 3-position is aryl substituted, but negatively impacts potency when the 3 position is alkylated. We therefore decided that rather than follow a rigid systematic investigation of each area of the molecule as an SAR follow-up strategy, we would use an empirical approach combining both single- and multi-dimensional SAR exploration. We made subtle variations around the 6- and 7-position substitution at the same time as exploring the 2- and 3-position. This was intended to give as broad a coverage of the chemotype as possible in a smaller number of compounds, and to avoid SAR elaboration of the core chemotype was focused on three missing key improvements masked by the hypothesized non-additive SAR.

**Synthesis and analoging**

Analoging was driven using the representative synthetic pathway shown in scheme 1. Various substituted 2-aminopyridines A were converted into 2-aryl imidazo[1,2-a]pyridines B via iodine-mediated condensation with arylmethylketones. Subsequent regioselective bromination of the 3 position of the imidazo[1,2-a]pyridines yielded 3-bromoimidazo[1,2-a]pyridines C, which were then converted into representative final products D via palladium mediated coupling with various primary and secondary amines. In some instances, representative final compounds
D were obtained directly in one step from aminopyridines A via tosic acid-mediated three-component cyclization with isonitriles and aryl aldehydes. One such example of D, 2-(pyridin-2-yl)-N-(2,4,4-trimethylpentan-2-yl)imidazo[1,2-a]pyridin-3-amine (E), was cleaved under acidic conditions to give 3-aminoimidazo[1,2-a]pyridine F, and subsequently converted into representative final compounds G via palladium mediated coupling with various aryl bromides. Further exploitation of intermediate F was achieved via amidation followed by borane mediated reduction of the amide to give N-alkylated representative final compounds K.

Structure activity relationship (SAR)

SAR elaboration of the core chemotype was focused on three distinct regions (tables 2-4). Scanning of the 3-position of the imidazo[1,2-a]pyridine core revealed some initial SAR of this position (table 2), suggesting that compounds with both amino alkyl and amino aryl moieties at this position could demonstrate anti-parasitic activity. Although small lipophilic alkyl groups demonstrated moderate anti-parasitic activities and selectivity indices (9, 12), presence of both basic (10, 15) and non-basic (13, 16) heteroatoms appear detrimental to anti-parasitic activity. Conversely, presence of additional polar heteroatoms or halogens in the case of aniline-type substitution in the 3-position seemed to enhance potency and in certain cases selectivity over host cell cytotoxicity (5, 21), although, as previously mentioned, this selectivity seemed to partly depend on the nature of additional substitution at positions 6 and 7 on the imidazo[1,2-a]pyridine core (22-25, 29-30). A combination of alkyl and aryl substituents on the 3-position amine appeared to result in an interesting combination of both potency and selectivity over background cytotoxicity (20). It became clear that there were extensive opportunities to modulate physicochemical properties of the compound at the 3-position of the chemotype.

Investigation of the 2-position of the core (table 3) revealed that small substitutions such as chloro and methoxy on the 4- and 5-position 2-pyridyl were tolerated (32, 33, 42-47) and could be exploited to tune microsomal stability. Moving the position of the pyridyl nitrogen to the 3 or 4 position removed anti-leishmanial activity (34, 35), however replacement of the 2-pyridyl with 5-membered nitrogen-containing heteroaromatic rings was tolerated in certain instances (51), and this change also had a major impact on the metabolic stability of the compound. Interestingly, replacement of the pyridyl with electron enriched phenyls such as 4-anilino (37,38) and 4-alkoxy substituted (44, 45) retained moderate activity and generally removed any evidence of cytotoxicity, suggesting that the anti-parasitic efficacy of the chemotype is likely not driven by a bidentate metal chelation effect. This observation of multiple non-2-pyridyl analogues further supported our earlier hypothesis regarding the redundancy of the PAINS flag for the initial hit 1.

Broader substitution of the 6- and 7-positions of the core imidazo[1,2-a]pyridine revealed a clear negative effect of withdrawing electron density from the core heterocycle, with cyano (54, 58) and trifluoromethyl (55, 59) substituents negatively impacting the antiparasitic effect (table 4). Attempts to probe further away from the core negatively impacted potency and/or cytotoxicity (62-65), however incorporation of heteroatoms in this position
could be used to improve microsomal stability (64). Scanning the impact of 6- and 7- substitution relative to the functionality at position 3 reconfirms the previous observation that SAR appears non-additive, with changes having negative or positive impact on potency and cytotoxicity depending on the nature of the 3-position substitution.

ADME and pharmacokinetic profiling

To complete the initial scoping of this chemical series, we profiled a selection of compounds in preliminary in vitro solubility and metabolic stability assays (table 5). In general compounds appeared poorly soluble at pH 7.4, although this could be optimized through incorporation of additional polarity in the 3-position of the core (25). Kinetic solubility at pH 2.0 was consistently favourable, and the compounds generally demonstrated rapid intrinsic clearance in both human and mouse microsomes, with the intrinsic clearance of both species tracking when moderate improvements were observed in microsomal stability for compounds 25 and 57, which again suggested that ADME property optimization was strongly tied to the presence of polar heteroatoms in the 3 position of the imidazo[1,2-a]pyridine. Compiling the broader SAR around intrinsic microsomal mouse clearance in vitro (Tables 2-5) shows that improvement was generally achievable through the introduction of additional polarity, for example via incorporation of additional nitrogen in the pendant aryl groups at positions 2- and 3- of the core, or through ether groups at the 7-position of the core scaffold. To understand the translation of these in vitro ADME parameters into in vivo exposure, compound 23 was profiled in mouse pharmacokinetic studies at 1 mg/kg i.v. and 50 mg/kg p.o. (Fig 3). Despite moderate in vitro CLint and in vivo bioavailability, 23 demonstrated high clearance in the mouse, resulting in relatively low overall exposure in terms of CMax and time of exposure after oral dosing, with no compound observed in the plasma after 24 hours. Such exposure relative to the anti-parasitic activity of the compound was considered insufficient to progress the compound to an in vivo infection model of Leishmania.

Conclusions

Work continues to optimize the imidazo[1,2-a]pyridine series informed by the SAR generated by the in-silico booster process and subsequent informed design. The primary focus for this future work will be further improvement of the in vitro ADME parameters in order to achieve prolonged exposure in vivo in line with a profile capable of demonstrating efficacy in an infection model of Leishmania. This may, however, be complicated by the clearly non-additive anti-parasitic and cytotoxicity SAR, although certain key findings from the SAR identified so far should enable improvement of anti-parasitic activity, microsomal stability and widening of the selectivity index between anti-parasitic effect and cell cytotoxicity. The overall approach of using the Drug Discovery Booster as a multi-party in silico screening approach to complete the first phase of hit-to-lead and hit validation allowed us to rapidly confirm and validate the activity and relevance of the series as well as inform and guide the initial SAR exploration. Further analysis of the impact and utility of the Booster approach, which has to date been applied to over 20 different hit series, will be the subject of future communication.
Conflicts of interest

There are no conflicts to declare.

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Notes and references

§ Compound libraries for HTS efforts at DNDi were sourced from a combination of commercial and proprietary collections.

§§ Assignment of blinded names A-E is random. Specific contributions of each partner were, and remain, anonymous within the collaboration.

§§§ Full synthetic details for all compounds in Supporting Information

§§§§ A full data set containing the compounds synthesized during this effort can be found in the supporting information and has been made fully available online: https://dndi.org/research-development/portfolio/drug-discovery-booster/

§§§§§ 3 was followed up further by subsequent rounds of in silico screening via this consortium approach, results to be communicated in future correspondence.

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Table 1: Overview of the in silico approach taken by each partner company, along with the number of compounds furnished from each partner.

| Company and computational approach | Number of compounds (actives)* |
|-----------------------------------|--------------------------------|
|                                   | On scaffold* | Scaffold hop* |
| A- ECFP4 similarity (initial Tanimoto cut-off 0.7 descending incrementally until sufficient compounds had been identified) | 45 (12) | 15 (3) |
| B- Series of substructure-based queries prioritized with Tanimoto similarity calculated by use of Morgan fingerprint approach. Final selection cherry picked by eye. | 69 (10) | 121 (18) |
| C- ECFP4 similarity (Tanimoto cut-off 0.6) and in-house fingerprint search (Tanimoto cut-off 0.7). Subsequent in-house "quality", commercial availability, and IP filters. | 85 (11) | 16 (0) |
| D- Daylight and ChemAxon fingerprint similarity ranking\(^{41,42}\) followed by Openeye pharmacophore alignment\(^{42,43}\) | 58 (12) | 56 (4) |
| E- The top scoring 150 compounds were selected by Tanimoto similarity calculation using the FCFP4 fingerprint, followed by refinement to 96 compounds based on maximized diversity. | 106 (11) | 86 (9) |
| All companies | 363 (56) | 294 (34) |

a) Actives defined as compounds with SI > 5 against one or both parasites; b) compounds with a 2-substituted imidazo[1,2-a]pyridine core; c) Compounds without a 2-substituted imidazo[1,2-a]pyridine core
Table 2 SAR exploration of the 3 position for *L. donovani*

| Compound | R3                  | R6 | R7 | IC₅₀ µM ᵃ | CC₅₀ µM ᵇ | Clₙᶜ ᶜ |
|----------|---------------------|----|----|-----------|-----------|--------|
| 4        | N-cyclopentylamino- | H  | H  | 1.80      | 42.5      |        |
| 5        | N-[2,3-dihydro-1,4-benzodioxin-6-yl]-amino | H  | H  | 1.16      | 25.6      |        |
| 9        | N-2,2,2-trifluoroethylamino- | H  | H  | 10.2      | >50       |        |
| 10       | N-2-(N,N-dimethyl)-ethyl-1,2 diamino- | H  | H  | >50       | >50       |        |
| 11       | N-2-methyl-2-hydroxy-propylamino- | H  | H  | 30.4      | >50       | 171    |
| 12       | N-cyclopropylmethylamino- | H  | H  | 3.57      | 19.4      |        |
| 13       | N-oxetan-2-ymethlamino- | H  | H  | >50       | >50       |        |
| 14       | N-tetrahydropyran-3-ylamino- | H  | H  | 29.4      | >50       |        |
| 15       | N- pyrolin-3-ylamino- | H  | H  | >50       | >50       |        |
| 16       | N-morpholin-         | H  | H  | >50       | >50       |        |
| 17       | N-benzyamino-        | H  | H  | 9.34      | >50       |        |
| 18       | N-phenylamino-       | H  | H  | 1.11      | 6.69      |        |
| 19       | N-4-fluorophenylamino- | Me | H  | 2.71      | 9.61      |        |
| 20       | N-(N-methyl)-4-fluorophenylamino- | H  | H  | 3.9       | >50       | >1000  |
| 21       | N-3-fluorophenylamino- | H  | H  | 3.70      | >50       | 242    |
| 22       | N-2-fluorophenylamino- | H  | OMe| 1.73      | 5.75      | 133    |
| 23       | N-3,5-difluorophenylamino- | H  | OMe| 1.91      | 4.3       | 62     |
| 24       | N-2,5-dichlorophenylamino- | H  | OMe| 3.91      | 8.93      |        |
| 25       | N-5-fluoropyridin-2-ylamino- | H  | OMe| 1.71      | 16.0      | 81     |
| 26       | N-pyrindin-2-ylaminoamino- | H  | Me | 4.53      | >50       | 130    |
| 27       | 4-trifluoromethoxyphenylamino- | H  | H  | 3.49      | >50       | 223    |
| 28       | N-4-cyanophenylamino- | H  | H  | 7.76      | >50       |        |
| 29       | N-4-methoxyphenylamino- | Me | H  | 2.31      | 10.7      |        |
| 30       | N-3-methoxyphenylamino- | Me | H  | 2.53      | 13.2      |        |
| 31       | N-2-methoxyphenylamino- | H  | H  | 2.61      | >50       | 326    |

ᵃ) *L. donovani* parasite clearance in infected THP1 cells; b) THP1 cytotoxicity; c) mouse microsomes, µL/min/mg pr
### Table 3 SAR exploration of the R2 position

| Compound | R3         | R2       | R6 | R7 | IC_{50} µM | CC_{50} µM | Cl_{int} µL/min/mg.pr |
|----------|------------|----------|----|----|------------|------------|------------------------|
| 4        | pyrid-2-yl | H H      |    |    | 1.80       | 42.5       |                        |
| 32       | 5-methoxy pyrid-2-yl | H H |    |    | 5.26       | 19.3       |                        |
| 33       | 4-methoxy pyrid-2-yl | H H |    |    | 1.76       | 27.1       |                        |
| 34       | pyrid-3-yl | H H      | >50|    |            |            |                        |
| 35       | 2-methyl pyrid-4-yl | H H |    |    | 42.2       | >50         |                        |
| 36       | 3,4-dimethoxy phenyl | H H |    |    | 16.2       | >50         |                        |
| 37       | Cyclohexyl | 4-(dimethylamino)phenyl | H Me |    | 7.37 (3.2) | >50 (29.3) | >1000                  |
| 38       | 4-methoxyphenyl | H Me |    |    | 7.9        | >50         |                        |
| 39       | 4-methoxyphenyl | H Me |    |    | 10.2       | >50         |                        |
| 40       | 2-fluoropyrid | H H |    |    | 25.9       | >50         |                        |
| 41       | 2-fluoro-4-methoxyphenyl | H H |    |    | 22.0       | >50         |                        |
| 42       | thiazol-2-yl | H H |    | >50|            |            |                        |
| 43       | N-methylimidazo2-yl | H H |    | >50|            |            |                        |
| 8        | 4-N,N-dimethylaminophenyl | Me H |    |    | 5.93       | >50         |                        |
| 44       | Cyclopentyl | 4-methoxyphenyl | H Me |    | 6.51       | >50         |                        |
| 45       | 4-ethoxyphenyl | H H |    |    | 9.4        | >50         |                        |
| 46       | 4-tertbutylyphenyl | H H |    |    | 13.2       | >50         |                        |
| 47       | 5-methoxy pyrid-2-yl | H H |    |    | 2.5        | 15.4       | 126                    |
| 48       | 4-fluorophenyl | H H |    |    | 2.85       | 4.22       | >1000                  |
| 49       | 5-chloro pyrid-2-yl | H H |    |    | 11.5       | >50         |                        |
| 50       | N-methylimidazo2-yl | H H |    | >50|            |            |                        |
| 51       | N-methylimidazo4-yl | H OMe |    |    | 4.46       | 40          | <12                    |
| 52       | 3-fluorophenyl | thiazol-4-yl | H OMe |    | 15.7       | 40.6        |                        |
| 53       | N-methyl pyrazol-3-yl | H OMe |    |    | 36.2       | >50         |                        |

**a)** L. donovani parasite clearance in infected THP1 cells; **b)** THP1 cytotoxicity; **c)** mouse microsomes, µL/min/mg.pr; **d)** value for library batch (value for resynthesized batch).
Table 4 SAR exploration of the 6 and 7 position of the core

| Compound | R3       | R6 | R7 | IC_{50} µM | CC_{50} µM | Cl_{in} |
|----------|----------|----|----|------------|------------|---------|
| 19       | Me       | H  | H  | 2.71       | 9.61       |         |
| 54       | CN       | H  | H  | >50        | >50        |         |
| 55       | CF3      | H  | H  | 3.54       | >50        |         |
| 56       | MeO      | H  | H  | 2.38       | 4.26       | 923     |
| 57       | Cl       | H  | H  | >50        | >50        |         |
| 58       | H        | CN | H  | >50        | >50        | 146     |
| 59       | H        | CF3| H  | 11.0       | 23.7       |         |
| 60       | H        | MeO| H  | 1.77       | 4.56       | 117     |
| 61       | H        | Me | H  | 0.71 (2.45) | 5.9 (11.8) | 355     |
| 21       | H        | H  | H  | 3.7        | >50        |         |
| 67       | H        | Me | H  | 2.47       | 14.24      |         |
| 62       | CH2OMe   | H  | H  | 7.19       | 16.0       |         |
| 63       | CH2NMe2  | H  | H  | 5.12       | >50        |         |
| 64       | H        | CH2OMe| H  | 2.74       | 7.37       | 92      |
| 65       | H        | CH2NMe2| H  | 4.54       | 33.3       |         |
| 66       | H        | OH | H  | 16.87      | 43.3       |         |
| 1        | Me       | H  | H  | 4.67 (2.9) | 28.0 (24.0)|         |
| 4        | H        | H  | H  | 1.80       | 42.5       |         |
| 68       | H        | Me | H  | 1.88       | >50        | >1000   |
| 5        | H        | H  | H  | 1.16       | 25.6       |         |
| 6        | Me       | H  | H  | 0.62 (1.98)| 16.0 (28.3)| 411     |
| 7        | Cl       | H  | H  | 1.91 (6.33)| >50 (17.4)| >1000   |
| 17       | H        | H  | H  | 9.34       | >50        | >1000   |
| 69       | Me       | H  | H  | 3.37       | >50        | >1000   |
| 18       | H        | H  | H  | 1.11       | 6.69       |         |
| 70       | H        | Me | H  | 0.82       | 9.59       |         |
| 10       | H        | H  | H  | >50        | >50        |         |
| 31       | H        | OMe| H  | 11.3       | >50        | 79      |
| 16       | H        | H  | H  | >50        | >50        | 137     |
| 71       | H        | OMe| H  | 17.9       | >50        |         |

a) *L. donovani* parasite clearance in infected THP1 cells; b) THP1 cytotoxicity; c) mouse microsomes, µL/min/mg.pr; d) library batch (*resynthesized batch*); e) library batch (*initial HTS batch*)
Table 5: in vitro ADME properties for select compounds

| Compound | Clint mL/min/mg pr | Kinetic solubility µg/mL |
|----------|--------------------|--------------------------|
|          | mouse  | human | pH 7.4 | pH 2.0 |
| 6        | 411    | 95    | 8       | 72     |
| 7        | >1000  | >100  | <0.5    | >76    |
| 25       | 81     | 35    | >76     | >76    |
| 37       | >1000  | >100  | <0.5    | 67     |
| 61       | 354    | >100  | <0.5    | 73     |
Figures

**Fig 1** Overview of Booster *in silico* screening process; First (A) and second (B) round of virtual screening result in data set of compounds with annotated SAR around 1; Data set is further annotated via expansive analoguing (C).
Fig 2 A) Diverse compounds identified from the Booster giving confidence in the viability of the chemical series; B) Dose response curves for 4; C) High content images of L. donovani infected THP1 cells treated with various concentrations of 4.
Fig 3 Pharmacokinetic profiling of 23 in mouse
Schemes

Scheme 1: i) ArCOCH₃, I₂, NaOH, H₂O, 110 °C; ii) NBS, MeCN, 20 °C; iii) R₃-NC, TsOH, ArCHO, MeOH, 70 °C; iv) R₃R₄NH, t-BuONa, Brettphos-Pd-G3, t-amylalcohol, 90 °C; v) HCl, MeOH, 20°C; vi) R₇COOH, EDCI, pyridine, 25°C; vii) BH₃SMe₂, THF, 60°C; viii) ArX, t-BuONa, Pd₂(dba)₃, XantPhos, PhMe, 110°C; ix) Ti(i-PrO)₄, NaBH₃CN, dioxane, 60°C