Synthesis of a Bicyclic Azetidine with In Vivo Antimalarial Activity Enabled by Stereospecific, Directed C(sp³)−H Arylation

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Supporting Information

ABSTRACT: The development of new antimalarial therapeutics is necessary to address the increasing resistance to current drugs. Bicyclic azetidines targeting Plasmodium falciparum phenylalanyl-tRNA synthetase comprise one promising new class of antimalarials, especially due to their activities against three stages of the parasite’s life cycle, but a lengthy synthetic route to these compounds may affect the feasibility of delivering new therapeutic agents within the cost constraints of antimalarial drugs. Here, we report an efficient synthesis of antimalarial compound BRD3914 (EC₅₀ = 15 nM) that hinges on a Pd-catalyzed, directed C(sp³)−H arylation of azetidines at the C3 position. This newly developed protocol exhibits a broad substrate scope and provides access to valuable, stereochemically defined building blocks. BRD3914 was evaluated in P. falciparum-infected mice, providing a cure after four oral doses.

INTRODUCTION

Malaria is caused by infection with Plasmodium species and transmitted by female Anopheles mosquitoes.1 While substantial progress has been made over the past two decades in reducing both mortality and burden associated with the disease,2 the emergence of parasite resistance to first-line antimalarial treatments highlights the need for safe and effective new therapies.1,3 The lifecycle of the malaria parasite includes three distinct stages—the asexual blood stage, the liver stage, and the gametocyte blood stage—and next-generation antimalarial therapies should ideally target all three.4

We recently identified antimalarial compounds originating from diversity-oriented synthesis (DOS), highlighted by phenylalanyl-tRNA synthetase inhibitor BRD7929 (2S,3R,4R) (Figure 1).5 BRD7929 exhibits activity in all stages of the parasite life cycle and is curative in multiple in vivo mouse models. However, a lengthy synthetic route to bicyclic azetidine BRD7929 (21 steps) hampers the ability to evaluate analogs systematically in vivo. Analysis of stereochemistry-based structure−activity relationships (SAR) revealed that two of eight possible stereoisomers of parent compound BRD3444—(2S,3R,4R) and (2R,3R,4R)—were active against a multidrug-resistant strain of malaria (P. falciparum, strain Dd2). This observation suggested that substituents at one of the stereocenters (C2) were not necessary for activity. Indeed, BRD3914, an analog lacking substitution at C2, was found to retain substantial antimalarial activity in vitro (EC₅₀ = 13 nM, Dd2 strain).6

Production cost constitutes an important consideration in the development of antimalarial therapeutics.4 We recognized in BRD3914 an opportunity to achieve an efficient synthesis of this class of antimalarials that proceeds with high regio- and

Figure 1. (a) Structures and in vitro antiplasmodial activities of representative bicyclic azetidines. (b) Stereochemistry-based structure−activity relationships (SAR).

Received: July 5, 2017
Published: July 21, 2017

DOI: 10.1021/jacs.7b06994
J. Am. Chem. Soc. 2017, 139, 11300−11306

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stereoselectivity of key transformations in fewer steps than the previous synthesis, and that minimizes the number of chromatographic purifications, which are known to drive both cost and waste output.\(^7\) To this end, we developed a stereospecific C(sp\(^3\))−H arylation of azetidines, enabling the preparation of all four stereoisomers of members of this series. The (2R,3S) stereoisomer of this building block led to an expeditious synthesis of antimalarial BRD3914. Subsequent in vivo evaluation in \(P. falciparum\)-infected humanized mice showed parasite clearance and lack of recrudescence after four oral doses, resulting in a durable cure.

A key step in our original synthesis of BRD3914 (14 steps) and other bicyclic azetidines comprises a base-mediated intramolecular cyclization from \(1\) that affords azetidine-2-carbonitrile \(2\) with little or no diastereoselectivity (ca. 1:1, Figure 2a).\(^6\) Following azetidine ring formation, nitrile reduction with DIBAL, protection with \(-\text{nitrobenzene-sulfonyl chloride, and allylation gives the precursor for the key ring-closing metathesis to generate the eight-membered ring (Figure S1). While the previous DOS route provided a versatile starting point for an initial SAR assessment focused, for example, on the 4-methoxyurea region, it limited the ability to assess other regions of the scaffold. Notably, various aryl and heteroaryl derivatives at the C3 position of the azetidine require the execution of the entire 14-step synthesis, and the introduction of functional groups on the eight-membered ring relies on olefin chemistry and thus presents regio- and stereoselectivity challenges.

In a revised retrosynthetic analysis (Figure 2a), we reasoned that a bifunctional linker could be used in an advantageous way to form the eight-membered ring (diamine 5). This approach would rely on \(cis\)-substituted azetidine 4 as a precursor, which we envisioned arising from a directed C−H arylation using commercially available \(p\)-azetidine-2-carboxylic acid 3. Since the first examples exploiting 8-aminoquinoline as a directing group by Daugulis and co-workers,\(^10\) C(sp\(^3\))−H functionalization methods have grown\(^11\) and been applied to a number of scaffolds, including cyclopropanes,\(^12\) cyclobutanes,\(^13\) cyclopentanes,\(^14\) pyrrolidines,\(^15\) and piperidines.\(^16\) There are few reports of the use of azetidines as a substrate\(^17\) in C−H functionalization, and to our knowledge the C3 arylation of azetidines has not been described.

**RESULTS AND DISCUSSION**

Unfortunately, initial attempts to extend C(sp\(^3\))−H arylation reactions to azetidines using known conditions exploiting 8-aminoquinoline as a directing group\(^6\) resulted in low yield of desired product (Table S1). Following substantial optimization on a model system (Figure 2b, Table S2−S9), we found that an unconventional \(N\)-trifluoroacetamide (N-TFA) protecting group\(^20\) and the additive (BnO)\(_2\)PO\(_2\)H\(^20\) were critical to reaction efficiency. Under the optimized conditions—aryl iodide (3 equiv), Pd(OAc)\(_2\) (10 mol%), AgOAc (20 mol%), (BnO)\(_2\)PO\(_2\)H (20 mol%), DCE (1.0 M), 110 °C—model product 7a was formed in 72% yield as determined by \(^1\)H NMR with internal standard. The reaction exhibited optimal performance under \(N_2\) atmosphere, though it afforded product in 52–56% yield under air or O\(_2\).

![Figure 2](image-url)
N-TFA is known to undergo hydrolysis readily, releasing the corresponding amine in mild alkaline solutions, raising the possibility that the azetidine could be deprotected directly after the arylation. We were delighted to find that upon completion of the C–H arylation, the N-TFA group could be cleaved in the crude reaction mixture by the addition of 7.0 N NH₃ in MeOH to afford the free azetidine. The one-pot C–H arylation/deprotection sequence was performed on a range of aryl iodides to demonstrate the broad applicability of this method (Table 1). The reaction tolerated a variety of 3- and 4-substituted aryl iodides (8a–m), including both electron-donating and electron-withdrawing groups. Halogenated aryl-iodide derivatives performed well and provide sites for further diversification. Exceptions include 2-fluoro-iodobenzene and 2-methyliodobenzene, which gave 32% isolated product (8n) and no product (8o), respectively, likely due to steric congestion around a palladacycle intermediate as proposed in previous studies. Increased yields were obtained with 4-iodobiphenyl (8p) and 2-
iodonaphthalene (8q). Arylation with 2-iodothiophene (8s) proceeded in 80% yield while other heterocyclic iodides were found to react less efficiently (8t−v). Encouragingly, the homologous pyrrolidine (9) and piperidine (11) analogs yielded products in 95% and 72% isolated yields, respectively (Figure 3a), demonstrating the utility of these reaction conditions in the installation of C3 aryl groups on other cyclic amines.

Next, we explored reaction conditions to remove the directing group (Figure 3b) to maximize stereochemical diversity. This would enable access to broad, well-defined chemical space in only a few steps from commercial material. We chose aryl bromide 8f as the substrate for these diversification studies given the possibility for further elaboration of the aromatic ring. Global Boc installation on (±)-8f (i.e., the azetidine and amide nitrogens) followed by a LiOH/H2O2-mediated cleavage of the 8-aminoquinoline auxiliary provided cis-(+)-13 in 69% yield with retention of stereochemistry at the α-position. Alternatively, mono-Boc protection followed by treatment with NaOH/EtOH at 110 °C afforded trans-(−)-14 in 81% yield, thus achieving both removal of the auxiliary and epimerization at the α-carbon. These conditions were then applied to (−)-8f and provided (−)-13 and (+)-14 in 72% and 77% yield, respectively. All four stereoisomers can be accessed from (+)- and (−)-8f without the need for chromatographic purification. Thus, any one of the four stereoisomers could be prepared in three steps with the right choice of substrate (1- or 3-azetidine-2-carboxylic acid) and auxiliary removal conditions.

Having established conditions for the C–H arylation and removal of the directing group, we turned our attention to the synthesis of BRD3914. TFA-protection and installation of 8-aminoquinoline on 3-azetidine-2-carboxylic acid (+)-3 (10.0 g) afforded 26.3 g of (+)-6a in 82% yield and could be achieved in a single reaction vessel (Figure 4). The use of only 0.95 equiv of DIPEA appeared critical to avoid epimerization at the α-position during the HATU-mediated amide coupling. The key one-pot arylation and deprotection sequence was performed on a 10-g scale with 1-bromo-4-iodobenzene and furnished (−)-8f in 52% yield with both relative and absolute stereochemistry confirmed by X-ray crystallography. Of note, we found that the reaction performed equally well on 0.3, 3.0, and 30.1 mmol scale (100 mg to 10.0 g, Table S10), although incomplete TFA-deprotection under standard NH3 in MeOH conditions was observed on ≥3.0 mmol scale. In these cases, modified conditions using K2CO3 in 9:1 MeOH/H2O completely removed the protecting group. Directing group cleavage from (−)-8f with retention of cis-stereochemistry afforded gram quantities of the synthetically useful carboxylic acid building block (−)-13.

Figure 3. (a) C–H arylation method to access homologous series of C–H arylation products. (b) Varying conditions for directing group removal to access selectively cis and trans key building blocks; all compounds are isolated as single stereoisomers with dr > 20:1 and ee > 99%.
Next, deprotection of (−)-17 using 4.0 M HCl in dioxane followed by reductive amination with bifunctional linker 15 proceeded smoothly in a single flask, delivering (+)-16 in 71% yield. Fmoc was removed using SiliaBond Piperazine, and sequential intramolecular amide bond formation led to lactam (+)-16 (structure confirmed by X-ray crystallography) in 89% isolated yield in a single flask. The lactam reduction proved remarkably difficult as conventional reagents such as LiAlH4 or BH3·Me2S resulted in either competing protodebromination or formation of intractable adducts. Ultimately, we found that modified conditions reported by Reeves and colleagues5,25 using 4.0 M HCl in dioxane reduced the lactam; the resulting diamine (−)-13 was converted to (+)-16 in 57% yield. Finally, a Pd-catalyzed Heck alkynylation with phenylacetylene delivered BRD3914 in 24 h (92%).

The short synthesis permitted further evaluation of the biological activity of BRD3914. First, the compound was confirmed to be potent in vitro (EC50 = 15 nM) against P. falciparum parasites (Dd2 strain), in line with previously measured activity.6b Pleasingly, BRD3914 exhibited low cytotoxicity to human cell lines (A549 CC50 = 38 μM, HEPG2 CC50 > 50 μM, HepG2 CC50 > 50 μM). We then sought to evaluate the in vivo efficacy of this compound for the first time against blood-stage P. falciparum 3D7HLH/BRD (expressing firefly luciferase) in nonobese diabetic/severe combined immunodeficiency (NOD/SCID) IL2−/−/Il2r−/− mice engrafted with human erythrocytes (huRBC NSG). This mouse model has been shown to correlate well with human malaria challenge models.4,26 huRBC NSG mice were inoculated with parasites (−48 h) before administration with either 25 mg/kg or 50 mg/kg BRD3914 (single dose at 0 h, or multiple dosing at 0, 24, 48, and 72 h, formulated in 70% PEG300/30% solution of 5% dextrose/H2O)5 and monitored for 30 days (Figures 5, S2). Chloroquine was used as a positive control. After a single 50 mg/kg dose of BRD3914, a reduction in parasite-associated bioluminescence was observed in mice, but recrudescence occurred around day three. Gratifyingly, huRBC NSG mice treated with four q.d. (quaque die) doses (25 mg/kg or 50 mg/kg) of BRD3914 were parasite-free after 30 days based on bioluminescent imaging. These results are striking compared to chloroquine treatment (4 × 25 mg/kg), where recrudescence of infection was observed in one of two mice.

While BRD3914 has a dosing shortcoming in vivo when compared to BRD7929, overall the compound succeeds in curing mice under a traditional Peters’ test,7 retains in vitro activity against P. falciparum parasites, and, most importantly, exhibits a more attractive safety profile.5 In our in vivo studies, BRD3914 outperformed the antimalarial chloroquine, one of the most successful antimalarial agents to date.28
The biological studies presented herein were facilitated by the development of a Pd-catalyzed C(sp³)−H arylation strategy, which allowed the efficient preparation of BRD3914 in seven steps with five chromatographic separations. This route represents a significant improvement over previous syntheses of this class of inhibitors with respect to step count, chromatographic separations, and waste minimization. Notable features of the synthetic approach are the stereospecificity of the arylation and the controlled cleavage of the directing group, which affords any one of four C3-arylated azetidine-containing antimalarials and antimicrobials in addition to finding general use for the preparation of diverse fragments for chemical libraries or peptidomimetics.

**CONCLUSIONS**

In summary, we developed a Pd-catalyzed, directed C(sp³)−H arylation of azetidines, which exhibited a broad substrate scope and permitted the preparation of stereochemically defined, synthetically useful carboxylic acid building blocks. We used this method in the short synthesis of antimalarial compound BRD3914, which cured *P. falciparum* infection in a mouse model after four oral doses. Collectively, our results arise from the ability to explore stereochemical space, we envision that this method can be applied to the synthesis of other promising antimalarials and antimicrobials in addition to finding general use for the preparation of diverse fragments for chemical libraries or peptidomimetics.

**ACKNOWLEDGMENTS**

We thank the Broad Institute analytical team for high-resolution mass spectrometry data, Tom Clinciemaille (Broad Institute) for assistance with SFC, Drs. Peter Müller and Jonathan Becker (Massachusetts Institute of Technology) for X-ray crystallographic structural analysis, and Drs. Michael F. Mesleh and Jenna B. Yehl (Broad Institute) for assistance with NMR. This work was supported by the Bill and Melinda Gates Foundation (grant OPP1032518). S.L.S. is an investigator at the Howard Hughes Medical Institute. M.M. was supported by a fellowship from the National Science Foundation (DGE1144152) and from Harvard University’s Graduate Prize Fellowship. J.Z. was supported by a postdoctoral fellowship from the German Academic Exchange Service (DAAD). B.M. was supported by a postdoctoral fellowship from Harvard University. O.V. was supported by a postdoctoral fellowship from the Wenner-Gren Foundations.

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Figure 5. huRBC NSG mice were inoculated with *P. falciparum* (3D7/HLH/BRD) blood-stage parasites 48 h before treatment, and BRD3914 was administered as a single dose (25 mg/kg or 50 mg/kg) at 0 h, or multiple dosing at (25 mg/kg or 50 mg/kg) at 0, 24, 48, and 72 h (n = 2 for each group, this study was conducted once). Chloroquine (CQ) was used as a positive control. Infections were monitored using the *in vivo* imaging system (IVIS). Bioluminescent intensity was quantified from each mouse and plotted against time. The dots horizontal line represents the mean bioluminescence intensity level obtained from all the animals before the parasite inoculation.