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Short Note

2-Cyclopropyl-6-phenyl-2,3-dihydrothieno[3,2-d][1,3,2]-diazaborinin-4(1H)-one

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Abstract: As part of our ongoing scaffold hopping work on antimalarial 2-aminothieno[3,2-d]-pyrimidin-4-one scaffold, we explored the dihydrothieno[3,2-d][1,3,2]diazaborinin-4(1H)-one as a potential new antimalarial series. Using conditions found in the literature, we obtained 2-cyclopropyl-6-phenyl-2,3-dihydrothieno[3,2-d][1,3,2]diazaborinin-4(1H)-one with 93% yield through a simple treatment. It was then characterized by NMR (1H and 13C) and HRMS. Given the structure of this molecule, its aqueous stability was assessed to determine its suitability for biological tests. To our knowledge, this is the first dihydrothieno[3,2-d][1,3,2]diazaborinin-4(1H)-one described.

Keywords: boron; diazaborininone; thiophene; aqueous stability

1. Introduction

Malaria is the world’s most prevalent parasitic disease in terms of deaths, with an estimated 229 million cases and 409,000 victims according to the World Health Organization (WHO) in 2019 [1]. It is caused by parasites of the Plasmodium genus (mainly P. falciparum) and transmitted by the infectious bite of Anopheles mosquitoes. The first line treatment, artemisinin-based combination therapies (ACTs), combines an artemisinin derivative with another antimalarial compound. However, the reduction of both deaths and infections is threatened by multiple factors including the spread of parasite resistance to artemisinin and its derivatives [1]. This resistance is caused by mutations of the kelch13 protein [2] and is characterized clinically by delayed parasitic clearance potentially leading to treatment failure of ACTs [3]. Initially discovered in South-East Asia, where they have become widespread [3], kelch13 mutations are now found in Africa, where they have recently been linked to in vitro resistances against artemisinin derivatives [4,5]. Thus, discovery of new antimalarial compounds with novel mechanisms of action is a priority in the battle against malaria.

In 2015, Cohen et al. described compound M1 (Figure 1) [6], a 2-aminothieno[3,2-d]pyrimidin-4-one hydrochloride salt displaying an all-stage activity against P. falciparum associated with low toxicity, no mutagenicity and possessing an unknown mechanism of action. Many thienopyrimidine compounds are described as anticancerous compounds, mainly kinase inhibitors [7] and one thienopyrimidinone compound, TAK-931 (simurosertib, Cdc7 inhibitor, Figure 1) is currently in phase II clinical trial for the treatment of metastatic cancers [8].

As part of our ongoing medicinal chemistry work to improve M1 properties, a scaffold hopping strategy on the thienopyrimidine core was implemented for structure-activity relationships purposes. This led to the idea of replacing the pyrimidinone moiety by a [1,3,2]diazaborinin-4-one moiety, seeking to synthesize compounds in thieno[3,2-d][1,3,2]diazaborinin-4-one series and assess their activity in vitro against multi-resistant P. falciparum K1 strain (Scheme 1).
Boron-containing compounds possess various properties such as anti-cancerous and anti-infectious activities [9]. Boron in approved drugs (Figure 2) is found either as boronic acid (in bortezomib or ixazomib) or incorporated into a cycle, such as oxaborole or oxaborinine (like tavaborole or vaborbactam respectively).

2. Results

We used reaction conditions described by Davies et al., starting from anthranilamide and various potassium trifluoroborates to afford benzo[d][1,3,2]diazaborinin-4-ones in the presence of the boron trifluoride-ethylamine complex [10]. Since the tert-butylamine moiety could not be transposed onto a [1,3,2]diazaborinin-4-one cycle, a cyclopropyl moiety was chosen as replacement. We decided to begin by synthesizing compound 2 (Scheme 2), since the starting materials are commercially available. Starting from 3-amino-5-phenylthiophene-2-carboxamide 1 and potassium cyclopropyltrifluoroborate with 3 equivalents of boron trifluoride ethylamine complex, we synthesized compound 2 in a 1:1 mixture of toluene and CPME after one day at 80 °C (Scheme 2). Small adjustments to the conditions in Davies et al. [10], were made, slightly increasing the quantity of potassium trifluoroborate (1.2 equivalent here vs. 1.05) and the reaction time (24 h here vs. 16 h). An excellent yield of 93% was obtained with simple filtration as extraction/purification protocol to afford target compound 2.
3. Discussion

The next step was to assess the antiplasmodial activity in vitro against *P. falciparum* strain, but this raised concerns as to the potential aqueous stability of compound 2. Indeed, benzo[d][1,3,2]diazaborin-4-ones obtained from anthranilamide are reported to be a protecting group for boronic acids and able to generate air-stable boronic acid precursors [11–13]. Davies et al. reported relative stability in neutral or acid pH for their benzo[d][1,3,2]diazaborin-4-ones derivatives, but with very small amounts of D$_2$O (from 6 to 16 µL).

Therefore, a stability assay was performed on 2 (Figure 3): the compound was dissolved in DMSO-$d_6$, a small amount of D$_2$O was added and the stability of the compound was then monitored via $^1$H NMR (see Materials and Methods for complete procedure). Upon addition of D$_2$O, signal changes can be seen on the two singlets around 9 ppm on the green spectra. After 16 h, 2 is completely degraded (Scheme 3, Supplementary Materials Figures S1, S4 and S5). New signals correspond to the starting material 1 [14] and cyclopropyl boronic acid [15] resulting from the hydrolysis of 2.

![Figure 3. Evolution of the NMR signals of compound 3 before and after addition of D$_2$O.](image)

Scheme 3. Observed degradation of 2 under neutral aqueous conditions.

Like benzo[d][1,3,2]diazaborinin-4-ones, 2 was found to be completely air stable up to one year after its synthesis when stored in a closed hemolysis tube. It is also stable in DMSO-$d_6$ only up to 72 h after dissolution (Supplementary Materials Figures S6 and S7).

To the best of our knowledge, this is the first description of thieno[3,2-$d$][1,3,2]-diazaborinin-4(1H)-one. However, its incompatibility with aqueous media makes it inappropriate as a potential drug candidate.
4. Materials and Methods

Starting materials were purchased from Sigma-Aldrich (Saint Louis, MO, USA) or Fluorochem (Derbyshire, UK). NMR spectra were recorded on a Bruker Avance NEO 400 MHz NanoBay spectrometer at the “Faculté de Pharmacie” of Marseille. The residual proton signal of the deuterated solvent was used as an internal reference: DMSO-\(d_6\) \(\delta = 2.50\) ppm for \(^1H\) and 39.52 ppm for \(^13C\). Data for \(^1H\) NMR are reported as follows: chemical shifts (\(\delta\)) in parts per million (ppm), multiplicity (described as follows: s, singlet; d, doublet; m, multiplet), coupling constants (\(J\)) in Hertz (Hz) and integration. Data for \(^13C\) NMR are reported as follows: chemical shifts (\(\delta\)) in parts per million (ppm). HRMS (ESI) spectrum (ESI) was recorded on a SYNAPT G2 HDMS (Waters) at the “Faculté des Sciences” of Marseille (St Jérôme campus). Melting points were determined on a Köfler melting point apparatus (Wagner & MunzGmbH, München, Germany) and are uncorrected.

2-Cyclopropyl-6-phenyl-2,3-dihydrothieno[3,2-d][1,3,2]diazaborinin-4(1H)-one (2): Under nitrogen atmosphere, 3-aminomethylthiophene-2-carboxamide (100 mg, 0.46 mmol), potassium cyclopropyltrifluoroborate (81 mg, 0.55 mmol) and boron trifluoride ethylamine complex (155 mg, 1.37 mmol) were suspended in toluene and cyclopentyl methyl ether (1:1 \(v:v\), 0.3 mol L\(^{-1}\)). The mixture was stirred for 24 h at 80 °C. Upon cooling, water (10 mL) was added, the mixture was filtered and the precipitate was washed with water affording 1.3 g of the product \((115\) mg, 93% yield). \(^1H\) NMR (DMSO-\(d_6\), 400 MHz) \(\delta\) 9.05 (s, 1H), 9.01 (s, 1H), 7.69 (d, \(3J = 7.3\) Hz, 2H), 7.52-7.37 (m, 3H), 7.21 (s, 1H), 0.75-0.66 (m, 2H), 0.63-0.55 (m, 2H), 0.12-0.01 (m, 1H). \(^13C\) NMR (DMSO-\(d_6\), 100 MHz) \(\delta\) 162.8, 150.9, 148.4, 133.0, 129.3 (2C), 129.2, 125.8, 115.2, 113.6, 5.3 (2C), 4.4. HRMS (ESI) \(m/z\) calculated for C\(_{14}\)H\(_9\)N\(_2\)OSB \([M+H]^+\) 269.0917, found 269.0915. mp = 249–251 °C (degradation).

Procedure for the aqueous stability test: 9 mg of 2 were dissolved in 500 µL of DMSO-\(d_6\). At \(t = 0.50\) µL of D\(_2\)O (10:1 \(v:v\)) were added. NMR analysis was then carried out on the sample resulting in the green spectra of Figure 3 (\(t \approx 5\) min). After 16 h (room temperature at 20 °C), another NMR analysis was performed resulting in the blue spectra of Figure 3 (\(t \approx 16\) h).

Supplementary Materials: The following are available online. Figure S1. \(^1H\) NMR spectra of 2-cyclopropyl-6-phenyl-2,3-dihydrothieno[3,2-d][1,3,2]diazaborinin-4(1H)-one, Figure S2. \(^13C\) NMR spectra of 2-cyclopropyl-6-phenyl-2,3-dihydrothieno[3,2-d][1,3,2]diazaborinin-4(1H)-one, Figure S3. HRMS (ESI) spectra of 2-cyclopropyl-6-phenyl-2,3-dihydrothieno[3,2-d][1,3,2]diazaborinin-4(1H)-one, Figure S4. \(^1H\) NMR spectra of 2-cyclopropyl-6-phenyl-2,3-dihydrothieno[3,2-d][1,3,2]diazaborinin-4(1H)-one after 5 min in DMSO-\(d_6\)/D\(_2\)O mixture (10:1 \(v:v\)), Figure S5. \(^1H\) NMR spectra of 2-cyclopropyl-6-phenyl-2,3-dihydrothieno[3,2-d][1,3,2]diazaborinin-4(1H)-one after 16 h in DMSO-\(d_6\)/D\(_2\)O mixture (10:1 \(v:v\)), Figure S6. \(^1H\) NMR spectra of 2-cyclopropyl-6-phenyl-2,3-dihydrothieno[3,2-d][1,3,2]diazaborinin-4(1H)-one after one year of storage, Figure S7. \(^1H\) NMR spectra of 2-cyclopropyl-6-phenyl-2,3-dihydrothieno[3,2-d][1,3,2]diazaborinin-4(1H)-one after 3 days in DMSO-\(d_6\).

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