Discovery of 1,3-Diaminobenzenes as Selective Inhibitors of Platelet Activation at the PAR1 Receptor

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Discovery of 1,3-Diaminobenzenes as Selective Inhibitors of Platelet Activation at the PAR1 Receptor

Chris Dockendorff,* Omozuanvo Aisiku,† Lynn VerPlank,‡ James R. Dilks,‡ Daniel A. Smith,‡ Susanna F. Gunnink,‡ Louisa Dowal,‡ Joseph Negri,† Michelle Palmer,† Lawrence MacPherson,† Stuart L. Schreiber,†§ and Robert Flaumenhaft* † Chemical Biology Platform and Probe Development Center, Broad Institute of Harvard and MIT, 7 Cambridge Center, Cambridge, Massachusetts 02142, United States
‡ Division of Hemostasis and Thrombosis, Department of Medicine, Beth Israel Deaconess Medical Center, Harvard Medical School, Boston, Massachusetts 02215, United States
§ Howard Hughes Medical Institute, Broad Institute of Harvard and MIT, 7 Cambridge Center, Cambridge, Massachusetts 02142, United States

Supporting Information

ABSTRACT: A high-throughput screen of the NIH-MLSMR compound collection, along with a series of secondary assays to identify potential targets of hit compounds, previously identified a 1,3-diaminobenzene scaffold that targets protease-activated receptor 1 (PAR1). We now report additional structure–activity relationship (SAR) studies that delineate the requirements for activity at PAR1 and identify plasma-stable analogues with nanomolar inhibition of PAR1-mediated platelet activation. Compound 4 was declared as a probe (ML161) with the NIH Molecular Libraries Program. This compound inhibited platelet aggregation induced by a PAR1 peptide agonist or by thrombin but not by several other platelet agonists. Initial studies suggest that ML161 is an allosteric inhibitor of PAR1. These findings may be important for the discovery of antithrombotics with an improved safety profile.

KEYWORDS: platelet activation, PAR1 inhibitor, allosteric inhibitor, 1,3-diaminobenzene, ML161, MLPCN MLSMR

Antiplatelet agents are an important part of regimens for patients at risk for adverse cardiovascular events, decreasing the probability of such events by minimizing thrombus formation following rupture of atherosclerotic plaque.1 However, contemporary antiplatelet drugs are only partially effective as evidenced by the substantial recurrence rate of arterial thrombosis despite current therapy.2 Furthermore, the beneficial effect of antiplatelet agents is tempered by an increased risk of dangerous hemorrhagic complications.3 Many developmental programs focused on new drug targets on platelets have been initiated over the past decade. Prominent among these programs have been those identifying and testing compounds that block protease-activated receptor 1 (PAR1)-mediated platelet activation.

Several inhibitors of PAR1 have been developed. The most advanced in clinical trials is vorapaxar (SCH530348), which was developed from a lead identified by a radioligand binding approach using a high affinity Thrombin Receptor Agonist Peptide.4 Vorapaxar is a potent inhibitor of PAR1 but was associated with an increased risk of intracranial bleeding when used in combination with standard therapy in a phase III trial (TRA-CER).5 Atopaxar (E5555) is a second PAR1 inhibitor in advanced clinical trials. Atopaxar therapy is associated with anti-PAR1 activity ex vivo; however, its use was associated with elevation of liver transaminases and QTc prolongation at higher doses in phase II trials.6–8 Other small molecule PAR1 inhibitors have been described9–13 but have not been tested in clinical trials. Safety and efficacy issues with current antiplatelet therapies, including investigational PAR1 antagonists, highlight the need for antiplatelet agents that may act via alternative mechanisms.

We recently reported14 a high-throughput screen of the National Institute of Health Molecular Libraries Small Molecule Repository (NIH-MLSMR) small-molecule library (∼300000 compounds), undertaken to identify inhibitors of granule secretion and/or platelet activation.15,16 The primary screen measured adenosine triphosphate (ATP) secreted from dense granules following SFLLRN-induced activation of PAR1 using a luciferin/luciferase detection system. Several chemically tractable scaffolds were identified that inhibited dense granule secretion. Target identification studies with hits from this screen showed that compounds with a 1,3-diaminobenzene core act at PAR1. This paper describes structure–activity relationship (SAR) studies of this class of compounds with the NIH-MLSMR compound collection, along with a series of secondary assays to identify potential targets of hit compounds, previously identified a 1,3-diaminobenzene scaffold that targets protease-activated receptor 1 (PAR1). We now report additional structure–activity relationship (SAR) studies that delineate the requirements for activity at PAR1 and identify plasma-stable analogues with nanomolar inhibition of PAR1-mediated platelet activation. Compound 4 was declared as a probe (ML161) with the NIH Molecular Libraries Program. This compound inhibited platelet aggregation induced by a PAR1 peptide agonist or by thrombin but not by several other platelet agonists. Initial studies suggest that ML161 is an allosteric inhibitor of PAR1. These findings may be important for the discovery of antithrombotics with an improved safety profile.

KEYWORDS: platelet activation, PAR1 inhibitor, allosteric inhibitor, 1,3-diaminobenzene, ML161, MLPCN MLSMR
PAR1 inhibitors as well as preliminary studies supporting an allosteric mode of action. Several compounds were identified by high-throughput screening (HTS) with a 1,3-diamidobenzene core. Of these, compound 4 was selected as a starting point for SAR studies, as it showed acceptable potency in the primary assay measuring inhibition of dense granule release (IC_{50} = 1–10 μM). Importantly, it was inactive in a luciferase counterscreen, and it showed little or no inhibition of platelet activation stimulated through other receptors (Figure S1 in the Supporting Information). Inhibition of PAR1 platelet activation by compound 4 was readily reversible. In addition, the compound did not inhibit phosphodiesterase 3A (PDE3A), which is present within platelets and promotes activation by suppressing cAMP levels. Compound 4 also showed acceptable solubility in PBS (40 μM) and good inhibition in a standard assay with human platelets measuring SFLLRN-induced surface expression of P-selectin (Table 1), a transmembrane cell adhesion molecule that is sequestered in platelet α-granules and expressed on the platelet surface following activation.

SARs of the 1,3-diamidobenzene scaffold are described in Tables 1–4. Compounds were prepared according to representative Schemes 1 or 2 (see the Supporting Information for details).

Keeping the west side of compound 4 fixed, the aryl substituent R1 was investigated comprehensively (Table 1), and analogues were tested in the P-selectin assay. Maximal activities were observed with ortho substituents (compounds 2–12); substituents at the meta and para positions had neutral or negative effects on potency (compounds 13–26). The best results were observed with electron-withdrawing or neutral lipophilic ortho groups, with the bromide substituent of compound 4 proving optimal, giving an IC_{50} of 0.26 μM. Methyl (8) and ethyl (9) also showed good potencies, although a larger ortho substituent (phenyl, 10) was not tolerated. A select number of heterocycles (furan, 3-pyridyl, and 3-quinolinyl, 27–29) were poorly active or inactive. In an attempt to improve the potency, disubstituted analogues 30–36 were prepared, but all showed lower potencies than 4.

Concurrent with our explorations at the east end of the scaffold, the alkyl chain at the west end was investigated (Table 2). Optimal potency was observed with a 3-carbon chain (4). Some potency was retained with the 2-carbon chain (38), but a 4-carbon chain (39) failed to demonstrate inhibition. Replacement of the alkyl chain with a phenyl ring (40) decreased activity at the target. We expected that branched alkyl chains could provide compounds with improved plasma stability (PS) over 4, which showed moderate stability in human plasma (80% remaining after 5 h) but poor stability in mouse plasma (<2%). Mouse plasma stable compounds are required for study in a number of our in vivo disease models, so we attempted to address this liability of 4. We hypothesized that branched alkyl chains could give compounds more resistant to proteases and esterases, and in fact, 42 and 43 showed some improvement. Compound 42 had only moderate potency but acceptable mouse PS (83% after 5 h). Cyclopentane 43 had acceptable potency (IC_{50} = 0.52 μM) but poor mouse PS (26%). The lack of activity observed with 1,4-diamide 44 indicates that the 1,3-substitution pattern about the central ring is important.

In a search for more potent, plasma-stable compounds, we continued our investigations by varying the central 1,3-diamidobenzene ring (Table 3). Several heterocyclic compounds were prepared, but neither pyridyl, benzimidazole, nor oxazole analogues (45–50) showed any significant activity in our assay. Substituents at the 2- and 6-positions of the central

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*SAR at the East End of the 1,3-Diamidobenzene Scaffold*
ring were not tolerated (51 and 52), although alternative groups could be worth investigating. Two compounds with reverse amides were screened (53 and 54), and the compound with the reverse amide at the western position (54) maintained much of the activity of the parent compound (3).

To explore SARs further and to search for compounds with improved potencies and physicochemical and biological properties (especially PS), additional compounds were prepared and screened. These results are listed in Table 4, which includes plasma protein binding (PPB) and PS. Replacement of the western propylamide of 4 with a carbamate (55) or sulfonamides (56 and 57) gave compounds with decreased potencies. N-Methyl amides 58–60 were inactive, which suggests that both amides act as hydrogen bond donors with the target. The constrained analogues 61–64 were similarly inactive. Removal of the eastern carbonyl group of 4 was tolerated somewhat (65), but interestingly, much of the activity was retained with removal of the western carbonyl group in aniline 66.

In a final attempt to address the mouse PS liability of 4, we made additional branched alkyl analogues, building upon the moderate results of 42 and 43. The more highly branched \( \alpha,\alpha \)-dimethylamide 67 was very plasma stable but poorly active, but moving the branching point further from the amide carbonyl (68) gave a compound nearly equipotent to 4 (IC\(_{50} = 0.29 \mu M\)) with good human PS (90% after 5 h) and mouse PS sufficient for in vivo work (65% after 5 h), as well as adequate solubility for a probe compound (20 \( \mu M \) in PBS). The preparation of 4 is described in Scheme 1,21 and the synthesis of the mouse plasma-stable analogue 68 is depicted in Scheme 2. Prior to these more recent studies, compound 4 was formally nominated as a molecular probe (ML161) for the study of platelet activation.14

The selectivity at PAR1 is supported by studies involving various platelet activators in the presence of ML161. Washed human platelets were separately treated with the peptide AYPGKF (a PAR4 agonist), PMA (a protein kinase C activator), U46619 (a thromboxane receptor agonist), or collagen (an agonist of collagen receptors). ML161 did not inhibit these activators to any significant degree (Figure S1 in the Supporting Information). ML161 displayed dose-depend-
ent inhibition of thrombin-induced platelet activation, as measured by P-selectin expression (Figure S2 in the Supporting Information). Inhibition of PAR2, a widely distributed protease-activated receptor, was not evaluated since this receptor is not present on platelets. ML161 displayed selective inhibition of SFLLRN and thrombin-induced platelet aggregation, both of which operate via PAR1, and had no effect on AYPGKF, thromboxane, or ADP-induced platelet aggregation, which are all agonists at alternative platelet GPCRs (Figure S3 in the Supporting Information).

Continued work on the 1,3-diaminobenzene scaffold was inspired by our discovery that its mode of action at PAR1 may differ from reported PAR1 orthosteric inhibitors.22,23 Activation of platelets via PAR1 cleaved by thrombin24 leads to multiple downstream effects and observable phenotypic changes, including granule release and shape change characterized by extended pseudopodia. Blockade of PAR1 by orthosteric antagonists, such as those currently under clinical investigation,25 would be expected to inhibit all phenotypic changes. In contrast, 4 inhibits granule secretion, but not platelet shape change, as monitored in a SFLLRN-induced platelet aggregation assay.26 We hypothesize that it may be acting in an allosteric manner to inhibit select G-protein-coupled pathways mediated by PAR1. Our preliminary studies suggest that it inhibits Gqα, which is required for granule release, but not G12/13 signaling, which affects shape change. Additional evidence for a nonorthosteric inhibition mechanism was obtained by evaluating dose–response curves of SFLLRN-induced P-selectin expression in the presence of varying concentrations of 4 (Figure 1). Instead of a rightward shift of

![Figure 1. Dose–response curves of SFLLRN-induced P-selectin expression in the presence of varying concentrations of the PAR1 inhibitor 4.](https://dx.doi.org/10.1021/ml2002696)
such as 4 (ML161) may inhibit PAR1 in an allosteric fashion, which could enable the selective modulation of platelet activation pathways. Allosteric inhibition of PAR1 could provide saturable receptor binding and selective modulation of downstream G-protein signaling pathways. These pharmacological attributes may decrease the risk of life-threatening hemorrhage in the setting of anti-PAR1 therapy.26 Analogues of the 1,3-diaminobenzene scaffold will be important probes for evaluating this hypothesis.

**ASSOCIATED CONTENT**

 Supporting Information

 Additional assay results (Figures S1 to S3), assay protocols, and synthetic procedures and characterization data for 4, 66, and 68. This material is available free of charge via the Internet at http://pubs.acs.org.

**AUTHOR INFORMATION**

Corresponding Author

*Tel: 1-617-714-7460. E-mail: cjdocken@broadinstitute.org (C.D.). Tel: 1-617-735-4005. E-mail: rflaumen@bidmc.harvard.edu. (R.F.)

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Notes

The authors declare no competing financial interest.

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**ABBREVIATIONS**

ATP, adenosine triphosphate; HTS, high-throughput screening; IC50, concentration of inhibitor giving half-maximal activity; NIH, National Institute of Health; MLPCN, Molecular Libraries Probe Center Network; NIH-MLSMR, National Institute of Health Molecular Libraries Small Molecule Repository; PAR1, protease-activated receptor 1; PPB, plasma protein binding; PBS, phosphate-buffered saline; PS, plasma stability; SAR, structure–activity relationship

**REFERENCES**


(17) Briefly, compounds were incubated in platelet-rich plasma in 384-well plates at a concentration of 7.5 μM (for the primary screen) or at a dose response (for confirmation assays) for 30 min. Next, a mixture of the CellTiter-Glo (Promega) reagent (for ATP detection with a luciferase/luciferin system) and SFLRNR (EC50 for platelet activation ~5 μM) was used. Each plate was incubated for 15 min at 22 °C, and then luminescence resulting from ATP-driven luciferin production was measured by a plate reader. See ref 13 for full details.


(19) For analysis of P-selectin expression, 20 μL of gel-filtered platelets (0.5–1 × 108/mL) were incubated with the indicated agonists and subsequently stimulated with 5 μM SFLRNR (EC10 for platelet activation ~5 μM) for 10 min. Following stimulation, 10 μL of reaction mixture was transferred to 5 μL of PE-conjugated AC12.1 anti-P-selectin antibody. Phosphate-buffered saline (PBS; 500
μL) was added after a 20 min incubation at 37 °C, and the platelets were analyzed immediately by flow cytometry. See the Supporting Information for further details.


(21) At the present time, compound 4 (ML161) is also commercially available from ChemBridge Corp.


(24) The serine protease thrombin cleaves the N terminus of PAR1, which then acts as a tethered ligand to activate signaling through intramolecular ligation of the orthosteric binding site. SFLLRN peptide is derived from this tethered ligand and is an agonist of PAR1 when applied externally. For the initial discovery, see Vu, T.-K. H.; Hung, D. T.; Wheaton, V. I.; Coughlin, S. R. Molecular cloning of a functional thrombin receptor reveals a novel proteolytic mechanism of receptor activation. *Cell* 1991, 64, 1057−1068.


(26) Details will be disclosed in a full paper currently in preparation.
