

Pyrimidine-Based Tricyclic Molecules as Potent and Orally Efficacious Inhibitors of Wee1 Kinase

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Supporting Information

ABSTRACT: Aided by molecular modeling, compounds with a pyrimidine-based tricyclic scaffold were designed and confirmed to inhibit Wee1 kinase. Structure-activity studies identified key pharmacophores at the aminoaryl and halobenzene regions responsible for binding affinity with sub-nM

 K_i values. The potent inhibitors demonstrated sub- μM activities in both functional and mechanism-based cellular assays and also possessed desirable pharmacokinetic profiles. The lead molecule, 31, showed oral efficacy in potentiating the antiproliferative activity of irinotecan, a cytotoxic agent, in a NCI-H1299 mouse xenograft model.

KEYWORDS: Weel kinase inhibitors, antitumor

ells utilize G1 and G2/M checkpoints within cell cycle to ensure genomic integrity. The fact that cancer cells are often defective in the G1 checkpoint due to frequent p53 mutations has led to interest in developing anticancer agents that abrogate G2/M arrest. 1,2 One of the key players in this arena is the Weel kinase, an atypical tyrosine kinase that phosphorylates Cdk1 (also called Cdc2) on tyrosine 15 (Y15) resulting in functional inactivation.³ Cdk1 recruits Cyclin A and Cyclin B to initiate mitosis. Wee1 negatively regulates Cdk1 and enables repair of damaged DNA in G2 phase prior to mitosis. Therefore, inhibition of Wee1 forces tumor cells into premature mitosis, leading to mitotic catastrophe and cell death. 4,5 Since Wee1 is overexpressed in certain malignancies, 6-8 Wee1 inhibitors have the potential to sensitize p53 deficient tumors to DNA-damaging treatments.

Recently, a few classes of Wee1 inhibitors have been disclosed. Among them is a selective inhibitor, MK-1775 (1, 2-allyl-1-(6-(2-hydroxypropan-2-yl)pyridin-2-yl)-6-((4-(4methylpiperazin-1-yl)phenyl)amino)-1H-pyrazolo[3,4-d]pyrimidin-3(2H)-one). MK-1775 exhibited antitumor activity in various preclinical studies in potentiating chemo- and radiotherapy and is currently in phase I/II human clinical trials. 12 In addition, a different class of Weel inhibitors featuring an imino-dihydropyrimidinone pyrimidine core (2) has emerged in patent literature. 13 Computer-assisted evaluation of the Weel kinase binding site, using public X-ray structures of bound Weel inhibitors as starting points, 14,15 revealed a key region within the ATP binding pocket in Wee1 adjacent to the bicyclic core not utilized by inhibitors such as 2. This region is generally occupied by the pyridyl moiety of MK-1775 (1) and was deemed to be accessible from the ligand hinge region and therefore exploitable for drug design purposes. Taking advantage of this untapped region by 2, our de novo design and modeling studies suggested that molecules with a tricyclic core, as displayed in 3, would possess key elements for interacting with the active site of the Weel protein (see Figure 1A for more details), potentially resulting in improved Wee1 inhibition. In this letter, we demonstrate that not only do such molecules show potent activity in enzymatic and cellular assays but they also exhibit desirable pharmacokinetic properties as well as oral antitumor efficacy in a murine xenograft model.

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Received: July 2, 2014 Accepted: August 4, 2014 Published: August 6, 2014 The general synthesis of analogs presented in this letter is outlined in Scheme 1. The known amino pyrimidine 4 was

"Reagents and conditions: (i) NaH, 1-Cl-2-isocyanatobenzene, 98%; (ii) POCl₃, DIEA, 84%; (iii) 2,2-dimethoxyethanamine; (iv) HCl, 59% (2 steps); (v) mCPBA, 92%; (vi) amine.

10-25

treated with NaH and a halo-substituted isocyanatobenzene to afford the bicyclic intermediate $5.^{16}$ The pyrimidinedione moiety was chlorinated in the presence of POCl $_3$ and DIEA. An S $_{\rm N}$ Ar addition using 2,2-dimethoxyethanamine led to 7, which was subsequently cyclized under acid conditions. The methyl thioether group of 8 was oxidized by mCPBA, setting the stage for the final S $_{\rm N}$ Ar reaction with a desired amino substrate to provide $10-25.^{17}$ Compounds 26 to 31 were synthesized in the same fashion using the corresponding bis-halo isocyanatobenzene in the first step.

The binding affinity of all compounds, presented as K_i values, was assessed in a routine 6-point TR-FRET binding assay. A functional antiproliferative cell viability assay and a mechanismbased pCDK1 ELISA assay, both utilizing the NCI-H1299 cell line (human nonsmall cell lung carcinoma) to provide EC50 values, helped to access cellular potency of any potent binders. It was gratifying to find that the first analogue synthesized, 10, was active in the biochemical assay ($K_i = 2.3 \text{ nM}$) and also possessed reasonable cellular activity in both the cell viability and ELISA assays (EC₅₀ = 1.0 and 0.37 μ M, respectively). According to the model of compound 10 bound to the Weel active site (Figure 1A), one of the pyrimidine nitrogens and the tethered amino group form hydrogen-bond interactions to the backbone NH and backbone carbonyl O of residue Cys379, respectively (i.e., a two-hinge interaction motif). The model also predicted a third, nonhinge, H-bond interaction between the side-chain N of Wee1 gatekeeper residue Asn376 and the amide carbonyl O in compound 10. This third H-bond interaction was believed to be critical for good kinase

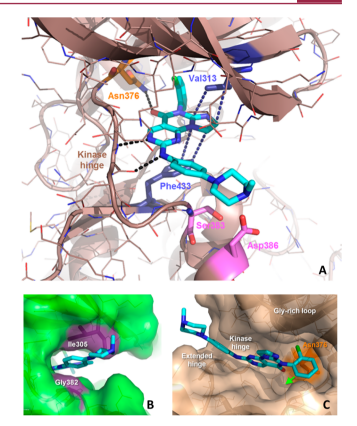


Figure 1. (A) Computational molecular model of compound 10 (shown in cyan, stick) bound to the Weel kinase binding site (shown in brown, ribbon). Key residues are shown in stick. The ligand is predicted to make three H-bond interactions with the protein (black dashes). Two of these interactions are made with backbone polar atoms in the hinge. The third interaction is made with the side chain of gatekeeper residue Asn376 (shown in orange, stick). The 5-membered ring added to the original bicyclic core (leading to the novel tricyclic core) is predicted to be sitting in a sweet spot right between Phe433 located at the base of the ATP binding site, and Val313 located in the Gly-rich loop. Purple dashes indicate the extent of the "sandwich" made by these two residues. (B) Close fit of benzene ring (part of R group) with surrounding protein walls, defined by residues Ile305 (Gly-rich loop) and Gly382 (extended hinge). (C) Halogenated phenyl ring uses small pocket next to gatekeeper residue Asn376 (shown in orange), while sitting perpendicular to tricyclic core. Green arrow indicates unutilized space.

selectivity since human kinases rarely have a hydrogen-bond donor (HBD) residue such as Asn in the gatekeeper position (<0.5% of ~500 human kinases based on internal studies).

Encouraged by these early results, we proceeded to map the structure—activity relationship (SAR) of the aminoaryl group attached to the pyrimidine ring. Removing the terminal piperazine on 10 led to about 10-fold loss in potency (11), while the importance of the aminoaryl moiety is clearly demonstrated with 12 being inactive ($K_i > 3140 \text{ nM}$) (Table 1). The reason for loss of activity by replacing the benzene with cyclohexane is likely due to the nature of the "gate" region leading to the ATP binding site (Figure 1B). Aromatic rings such as benzene are the ideal fit for this region in the protein, i.e., they are properly "sandwiched" between residue Gly382 (located in the extended hinge region) and residue Ile305 (located in the Gly-rich loop). Pi—sigma interactions between the aromatic ring in the ligand and aliphatic C—H bonds in the protein (alpha-C of Gly382 and beta-C of Ile305) are the main drivers behind the optimal

Table 1. Mono-Cl Analogues

		=1		
ID	R	Ki	H1299	pCDK1
		(nM)	EC ₅₀ (μΜ) ^a	$EC_{50} (\mu M)^b$
10	-\(\)_\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	2.3	1.0	0.37
11		21	5.9	6.8
12	$\vdash \!$	>3140	N.T.	N.T.
13		87	>10	>10
14	-<	5.5	9.8	3.8
15	-{	8.3	9.6	3.1
16	-\(\)\n_\n_\n	6.6	3.3	1.1
17	~~~~~	6.2	6.5	1.5
18	N_N_N-	>3140	N.T.	N.T.
19	VIII VIII VIII VIII VIII VIII VIII VII	26	7.3	6.0
20	√ N N	1.6	0.77	0.22
21	V	8.6	>10	2.3
22	V N	7.4	9.3	2.7
23	V N	1.2	0.69	0.10
24	V	5.4	2.0	0.62
25	V N	<1.0	0.55	0.098

^aCell viability assay. ^bELISA assay in H1299 cells. All data were an average of at least two measurements except for those with a qualifier or high micromolar values. N.T.: not tested.

protein—ligand complementarity observed in this region. Unlike benzene, saturated rings such as cyclohexane are too voluminous to fit well in this somewhat narrow region of the kinase binding site. At least one nitrogen atom of the piperazine on 10 was needed to maintain a K_i value below 10 nM (13–15), although reducing the basicity of the terminal nitrogen through acylation had little impact on Wee1 binding (16). Highly hydrophobic groups such as the cyclohexyl on 13 do not match well with the local environment in this solvent exposed region of the kinase, dominated by polar residues Ser383 (ribose residue) and Asp386 (ribose + 3 residue) shown

in Figure 1A. Once the ligand tail starts to project into solvent, R groups with a larger polar surface area (PSA), such as those in 14-16, are much better choices. Moving the piperazine to the meta position of the benzene maintained the potency (17, $K_i = 6.2 \text{ nM}$); however, placing it at the ortho position was not tolerated (18). Piperazine at the meta position confers the ligand a global conformation that still matches the V-shape/Cshape needed to complement the kinase binding site. Unfortunately, piperazine at the ortho position locks the ligand in a new conformation that is no longer compatible with the shape of the target binding site. Next, we investigated the SAR of bicyclic scaffolds with a saturated ring fused to the benzene. Once again, the presence of a nitrogen atom on the saturated ring was desirable for enzymatic potency (19 vs 20-22). Among these analogues, the tetrahydroisoquinoline inhibitor, 20, exhibited much better cellular activities and was the focus of further optimization. Installing a cyclopropyl group at the 4position of the isoquinoline of 20 led to 23, another analogue not only with high enzymatic potency ($K_i = 1.2 \text{ nM}$) but also possessing sub-micromolar activity in the cell viability (EC₅₀ = 0.69 μ M) and pCDK1 ELISA (EC₅₀ = 0.10 μ M) assays. Acylation of the tetrahydroisoquinoline nitrogen brought a slight deterioration in activity (24). The gem-dimethyl analogue, 25, showed overall potent activity comparable to 23. Four representative compounds (10, 20, 23, and 25) were evaluated for their pharmacokinetic (PK) profile in mice (Table 2). Even though they all demonstrated desirable profiles, 25

Table 2. Mouse PK for Mono-Cl Analogues^a

ID	$iv t_{1/2}$ (h)	iv CL (L/h/kg)	po AUC $(\mu M \cdot h)$	po F (%)
10	1.9	1.8	8.4	72
20	1.5	2.2	5.2	52
23	2.3	1.5	9.8	70
25	2.8	1.4	12.3	80

^aCD-1 mouse. iv dosing: 3 mg/kg. po dosing: 10 mg/kg.

appeared to have the best overall properties. The iv half-life reached 2.8 h, while clearance was modest (1.4 L/h/kg) with oral biovailability of 80%.

Having gained much insight in the SAR of the aminoaryl region, we turned our attention to the modification of the chloro-benzene ring (Table 3). Modeling suggested that the ortho-Cl group is essential for situating the phenyl moiety orthogonal to the tricyclic core (Figure 1C). Not only is the intramolecular arrangement optimal when the phenyl group is rotated almost 90° with respect to the plane of the tricyclic core, but the space available in the pocket (adjacent to gatekeeper residue Asn376) is also limited, and it can only accommodate such a large ring if placed "facing" the gatekeeper residue. The benefit of adding an extra halo substitution at the other ortho position is 2-fold. On one hand, the 2,6disubstitution pattern reinforces the 90° rotation needed for optimal complementarity with the immediate surroundings. On the other hand, the second halogen fills the remaining portion of the pocket (otherwise empty, i.e., with only a monosubstituted phenyl), increasing van der Waals contacts with the protein, and further increasing compound affinity/potency. Not surprisingly, all bis-halo analogues (26 to 31) demonstrated potent activity in the biochemical assay with K_i values below the lower detection limit for all except 27. The potent binding affinity was presumably responsible for the generally superior cellular activities possessed by this group of compounds. Four

Table 3. Bis-Halo Analogues

ID	R	X	Ki (nM)	H1299 EC ₅₀ (μΜ) ^a	pCDK1 EC ₅₀ (μM) ^a
26	-/_N_N_N-	Cl	<1.0	0.19	0.087
27		F	2.0	0.45	0.17
28	V N	Cl	<1.0	0.22	0.027
29	V	F	<1.0	0.21	0.029
30	V	Cl	<1.0	0.23	<0.013 ^b
31	V	F	<1.0	0.31	0.060

^aAverage of two measurements. ^bValues of 0.013 and <0.010 μ M.

compounds (26 and 29–31) were chosen for mouse PK studies with comparably favorable properties to that observed for the mono-Cl compounds (Table 4). The best compound, 31, achieved an adequate $i\nu$ half-life (2.8 h), low clearance (0.55 L/h/kg), and excellent oral biovailability (F = 94%).

Table 4. Mouse PK for Bis-Halo Analogues^a

ID	$iv \ t_{1/2} \ (h)$	iv CL (L/h/kg)	po AUC (μM·h)	po F (%)
26	1.7	2.0	6.5	55
29	2.6	1.1	12	68
30	3.9	0.97	9.8	49
31	2.8	0.55	24	94
^a CD-1 mouse. iv dosing: 3 mg/kg. po dosing: 10 mg/kg.				

Compound 31 was dosed orally at 100 mg/kg/day (mkd) in H1299 tumor bearing mice (n=4), utilizing inhibition of pY15 Cdk1/2 as an *in vivo* pharmacodynamics marker of Wee1 inhibition. The resulting Western immunoblotting analysis (Figure 2) demonstrated a nearly complete depletion of the pY15 bands at both 6 and 10 h after dosing. The concentration of 31 in plasma decreased 32% between these two time points, however, the tumor exposure levels remained at high throughout (47 and 53 μ M).

Compound **31** was further evaluated for its activity in potentiating the efficacy of the cytotoxic agent, irinotecan, in a mouse xenograft model (H1299). As presented in Chart 1, **31** achieved a modest tumor growth inhibition (TGI) of 43% (p < 0.001) after oral dosing at 50 mkd on day 41. Meanwhile, irinotecan alone reached a TGI of 67% (p < 0.001) at 10 mkd. The combination of **31** and irinotecan achieved an 83% TGI (p < 0.001). On day 52, the combination group showed a 58% TGI when compared to irinotecan alone (p < 0.05). The durability of the combination response was highlighted by a 25% tumor growth delay (TGD) when compared to irinotecan

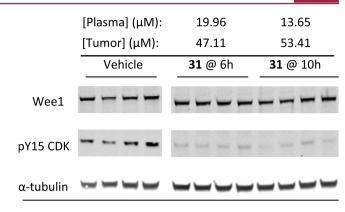
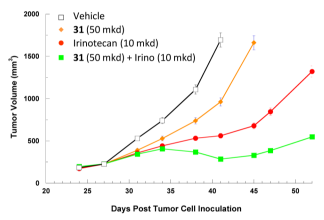


Figure 2. Pharmacodynamic profiles of compound **31**. Biomarker effects of **31** on expression of pY15 in H1299 tumor bearing mice after po dosing (100 mkd) as analyzed by Western blot. Loading control: α -tubulin.

Chart 1. Tumor Growth Inhibition^a



"Mouse xenograft (H1299 cell) data with tumor volume 52 days posttumor inoculation. Compound 31 was orally dosed QD ×14 from day 28 to day 41. Irinotecan was dosed IP at days 28, 32, 36, and 40.

alone (p < 0.01). All dosing regimens were well tolerated. Furthermore, the efficacy of **31** to sensitize irinotecan appeared to be dose responsive.¹⁹ Together, these results highlight the combination efficacy of **31** with irinotecan *in vivo*.

As disclosed in this letter, compounds with a pyrimidine-based tricyclic core were suggested by design/modeling and supported by their binding and cellular activity as potent Wee1 inhibitors. These molecules are generally selective against other kinases. SAR studies in the aminoaryl and halo-benzene regions led to compounds with sub-nM K_i values in the enzymatic assay and sub- μ M EC₅₀ values in both functional and mechanism-based cell assays. Representative inhibitors also exhibited desirable pharmacokinetic profiles. Compound 31 demonstrated oral *in vivo* efficacy in a mouse H1299 xenograft model and potentiated the antiproliferative activity of irinotecan. These results suggest that this class of compounds may have potential as therapeutic agents for anticancer indications.

ASSOCIATED CONTENT

S Supporting Information

Assay protocols, kinome selectivity profile, modeling protocols, efficacy study protocols, and synthetic procedures. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

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- (17) Synthesis of 20 involved the addition of *tert*-butyl 7-amino-3,4-dihydroisoquinoline-2(1*H*)-carboxylate followed by Boc removal and methylation on the isoquinoline nitrogen. Synthesis of 24 involved the addition of *t*-butyl 7'-amino-1'*H*-spiro[cyclopropane-1,4'-isoquinoline]-2'(3'*H*)-carboxylate followed by Boc removal and acylation on the isoquinoline nitrogen. See Supporting Information for detailed procedures.
- (18) Kinome profiling of a few key inhibitors against 84 kinases is provided in the Supporting Information.
- (19) Efficacy of 31 at a higher dose and more detailed biology studies will be published elsewhere.