Article

Structural Basis of Beneficial Design for Effective Nicotinamide Phosphoribosyltransferase Inhibitors

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Abstract: Inhibition of nicotinamide phosphoribosyltransferase (NAMPT) is an attractive therapeutic strategy for targeting cancer metabolism. So far, many potent NAMPT inhibitors have been developed and shown to bind to two unique tunnel-shaped cavities existing adjacent to each active site of a NAMPT homodimer. However, cytotoxicities and resistances to NAMPT inhibitors have become apparent. Therefore, there remains an urgent need to develop effective and safe NAMPT inhibitors. Thus, we designed and synthesized two close structural analogues of NAMPT inhibitors, azaindole–piperidine (3a)- and azaindole–piperazine (3b)-motif compounds, which were modified from the well-known NAMPT inhibitor FK866 (1). Notably, 3a displayed considerably stronger enzyme inhibitory activity and cellular potency than did 3b and 1. The main reason for this phenomenon was revealed to be due to apparent electronic repulsion between the replaced nitrogen atom (N1) of piperazine in 3b and the N8 atom of His191 in NAMPT by our in silico binding mode analyses. Indeed, 3b had a lower binding affinity score than did 3a and 1, although these inhibitors took similar stable chair conformations in the tunnel region. Taken together, these observations indicate that the electrostatic enthalpy potential rather than entropy effects inside the tunnel cavity has a significant impact on the different binding affinity of 3a from that of 3b in the disparate enzymatic and cellular potencies. Thus, it is better to avoid or minimize interactions with His191 in designing further effective NAMPT inhibitors.

Keywords: nicotinamide phosphoribosyltransferase; NAD+ biosynthesis; inhibitor; azacyclohexane; anticancer drug; drug design; enthalpy effect

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1. Introduction

Nicotinamide adenine dinucleotide (NAD\(^+\)) is a critical molecule in control of numerous basic cellular processes, such as ATP production and maintenance of cellular integrity and genome stability [1–4]. It not only serves as a cofactor for redox bioreactions but can also be a substrate for multiple NAD\(^+\)-consuming enzymes, such as poly(ADP-ribose) polymerases, mono-ADP-ribosyltransferases and sirtuins, participating in the epigenetic regulation of DNA transaction (transcription, replication, repair and recombination), cellular signaling processes and calcium homeostasis, and thereby in cell proliferation, differentiation and death [3–8]. Given the biological importance of this molecule, mammalian cells have evolved multiple biosynthetic pathways to produce NAD\(^+\); it is synthesized from tryptophan, nicotinic acid and nicotinamide (NAM)/nicotinamide riboside via de novo, Preiss–Handler and salvage pathways, respectively [1–4,9–11].

As NAD\(^+\) is consumed through many enzymatic processes, accelerated NAD\(^+\) depletion is often characteristic in cancer cells [9–12]. To rapidly replenish the NAD\(^+\) pool, cancer cells rely heavily on the NAM salvage pathway that backs NAM to NAD\(^+\) in two steps primarily catalyzed by nicotinamide phosphoribosyltransferase (NAMPT) [13–17]. Furthermore, in various types of cancer cells, NAMPT is found to be up-regulated [8,13,16,17], although the molecular mechanisms that dictate the salvage pathway choice remain elusive. Thus, NAMPT is considered an attractive target for the development of new anticancer drugs and therapies [18–31].

NAMPT, which functions as a homodimer with two unique tunnel-shaped cavities existing adjacent to each active site at the dimer interface, catalyzes the rate-limiting primary step of the salvage pathway, the transfer of phosphoribosyl residue from 5-phosphoribosyl-1-pyrophosphate (PRPP) to NAM to produce nicotinamide mononucleotide (NMN) (Figure 1) [1–4,32–34]. Many NAMPT inhibitors, such as FK866 (1), CSH-828, GNE0617 and STF-11880, which bind to the tunnel cavity of NAMPT, have been reported to exert anti-tumor suppression effects and have progressed to clinical trials [18,23–25,30]. However, cytotoxicities dominated by gastrointestinal symptoms and thrombocytopenia and resistance to NAMPT inhibitors have become apparent [23,35–37]. Therefore, there remains an urgent need to develop effective and safe anticancer NAMPT inhibitors for cancer chemotherapy. Thus, detailed insights into the tunnel structure of NAMPT are required to generate effective NAMPT inhibitors with a better therapeutic index and that can overcome resistance.

In the present study, we designed and synthesized two close structural analogues of NAMPT inhibitors 3a and 3b, which were modified from the best explored NAMPT inhibitor 1 [18,35]. Interestingly, we showed that 3a has a considerably stronger enzyme inhibitory activity and cellular potency than does 3b or 1. Furthermore, using these inhibitors as chemical probes, we characterized the inhibitor-targeting tunnel cavity of NAMPT. Importantly, our in silico binding mode analyses of these inhibitors with the tunnel cavity of NAMPT revealed that there is apparent electronic repulsion only between the nitrogen atom of piperazine in 3b and the Nδ atom of His191 in NAMPT. These findings indicate that the electrostatic enthalpy potential inside the tunnel region has a significant impact on the different binding affinity of 3a from 3b in the disparate cellular potencies. Thus, these results provide new insights into the design for further effective NAMPT inhibitors for cancer chemotherapies.
Figure 1. Overview of the human (h) nicotinamide phosphoribosyltransferase (NAMPT) as the dimer form and its enzyme reaction. The homodimer structure of hNAMPT (PDB Code: 2GVJ) [38–42] is shown in the ribbon diagram colored light blue and brown, respectively. The two tunnel-shaped cavities near the active sites in the hNAMPT molecule are shown in magenta shading. NAMPT catalyzes the conversion of nicotinamide (NAM) and phosphoribosyl pyrophosphate (PRPP) to produce nicotinamide mononucleotide (NMN).

2. Results and Discussion

2.1. Synthesis and Biochemical Properties of 3a and 3b

Many NAMPT inhibitors bind to the tunnel-shaped cavity of NAMPT (Figure 1), although the role of this tunnel in NAMPT function remains unknown [18–31]. These inhibitors have a unique pharmacophore consisting of three parts (Figure 2): head (a pyridine-like aromatic moiety), linker (a tunnel-interacting moiety) and tail (a solvent-exposed bulky group). To get more valuable insights into the tunnel cavity for the development of effective NAMPT inhibitors, we structure-basically designed and synthesized two close structural analogues of NAMPT inhibitors, namely azaindole–piperidine (3a)- and azaindole–piperazine (3b)-motif compounds, which were modified from the head and linker moieties of 1 (Figure 2, Scheme 1). Since the tail acts as a packing moiety against the solvent exposed tunnel exit surface, the tail structures of 3a and 3b were fixed as the same benzoyl group of 1. The nitrogen-containing aromatic heads present in many NAMPT inhibitors mimic the natural NAM substrate of the enzyme [18–31]. However, the head moiety of vinylpyridine ring of 1 (Figure 2) is known to be able to non-specifically interact with proteins [43,44]. Based on this knowledge, we first replaced this head to the 5-azaindole heterocyclic ring to improve the specificity to NAMPT (Figure 2). Secondly, as the linker moiety of azacyclohexane may be considered to become a critical motif for NAMPT inhibitory activity, we synthesized a close structural set of piperidine (3a)- and piperazine (3b)-motif NAMPT inhibitors according to the methods of Bair et al. [45] and Vogel et al. [46], respectively (Scheme 1).
Figure 2. Chemical structures of 3a, 3b and 1.

Scheme 1. Synthetic route of compounds 3a and 3b. Reagents and conditions: (i) NH₂NH₂, EtOH, r.t. to reflux, 2 h; (ii) 1H-pyrro[3,2-c]pyridine-2-carboxylic acid, EDC-HCl, DIPEA, DMF, r.t., 24 h. The targets 3a and 3b were prepared according to the methods of [45,46], respectively, with slight modifications.

To compare the NAMPT inhibitory activities of 3a, 3b and 1, we determined the IC₅₀ values by the titration curves from the standard calorimetric enzyme-cycling assay [31,36,37]. As shown in Figure 3a, the inhibitory activity of 3a (IC₅₀ = 0.11 μM) was approximately 5-fold stronger than that of 1 (IC₅₀ = 0.52 μM). This result suggests that the replacement of vinylpyridine with an azaindole ring could improve anti-NAMPT activity. The binding mode analyses of these inhibitors on the NAMPT molecule support this observation, as described in the next section; a hydrogen bond formed between the N1 atom of azaindole and Asp219 of NAMPT (Figure 4b). Surprisingly, the substitution of piperidine (3a) with piperazine (3b) resulted in a dramatic loss in inhibitory activity against NAMPT; the inhibitory activity of 3b (IC₅₀ = 5.06 μM) was shown to be about 50-fold weaker than that of 3a. The inhibition curve of 3a was much steeper than that of 3b, suggesting that incubation with 3a
resulted in the formation of an entity capable of extremely tight binding to the tunnel cavity of NAMPT. From these observations, we suspect that unfavorable interactions between the piperidine moiety of 3b and the tunnel cavity of NAMPT were responsible for the considerably weak anti-NAMPT activity.

**Figure 3.** Biochemical properties of 3a and 3b. (a) The inhibitory activities of 3a (open circle), 3b (open triangle) and 1 (open square) against NAMPT enzyme were examined, as described previously [31,36,37]. NAMPT activity was measured using a coupled-enzyme reaction system (CycLex NAMPT colorimetric assay kit) in 96-well plate format using the one-step method. Data are the averages of three independent experiments, and the bars indicate the standard error (SE) values. (b) The anti-proliferative effects of 3a (circle) and 3b (triangle) on HCT116 cells were investigated as described previously [31,36,37]. HCT116 cells were treated with the indicated concentrations of each inhibitor with (closed symbol) or without (open symbol) 100 µM NMN for 72 h. The cell activity (% of control) was measured by use of the WST-8 assay. Data are presented as the means of three independent experiments ± SE, and the bars indicate the SE values.

To confirm the marked differences of anti-NAMPT activities between 3a and 3b, the anti-proliferation effects of these inhibitors on the human colon cancer cell line HCT116 were examined after continuous exposure for 72 h [31,36,37]. As shown in Figure 3b, compound 3b displayed significantly weaker cell activity; the 50% effective concentration (EC50) value was calculated to be 87.5 nM, which was approximately 200-fold higher than that of 3a (EC50 = 0.48 nM). These results of the different inhibition degrees in cell-based assay from those in enzyme assay (Figure 3) paralleled observations by others that the degrees of potency of cell-based assessments were not always identical with those of NAMPT enzymatic inhibition [26,28,47]. In addition, there was little cytotoxicity on non-cancer cell lines, such as TIG-120, WI38 and MRC-5 cells. Both 3a and 3b caused more than 90% reduction of NAD+ levels in the cancer cells after 24 h-treatment and thereby induced cell death. Furthermore, supplementation of the product of NAMPT, NMN, to the cell culture medium could significantly rescue HCT116 cells from cell death by treatment with 3a or 3b (Figure 3b). Additionally, the dose–response curves of 1 were similar to those of 3a. These results confirmed that NAMPT is a target of both inhibitors. Collectively, these results indicate that the substitution of one carbon (C) atom of the piperidine moiety in 3a for a nitrogen (N) atom (piperazine moiety in 3b) exerts a critical effect on the anti-NAMPT activity.

### 2.2. Binding Modes of 3a and 3b to the Tunnel Cavity of NAMPT

To understand the different NAMPT inhibitory activities of 3a and 3b, we analyzed their binding modes to NAMPT (PDB Code: 2GVJ [38–42]) by our in silico binding mode analyses using RDKit [48] and LigandScout [49]. The superimpositions of these inhibitors in the tunnel cavity revealed that they adopted almost identical chair binding forms as with the case in 1 (Figure 4a). The NAMPT residues that contacted these inhibitors were observed in very similar locations in the structure. These results
indicate that the binding modes of 3a and 3b in the tunnel cavity of NAMPT do not significantly alter as compared with that of 1.

When observed closely, the position of the head regions of 3a and 3b in the tunnel cavity were nearly identical to that of 1 (Figure 4b). The subtle shifts observed in 3a and 3b from the position of 1 may have been introduced by their slightly different orientations of the head-azaindole bicyclic ring from the head-pyridine ring of 1 inside the tunnel cavity of NAMPT. The azaindole rings of 3a and 3b were sandwiched between Phe193A and Tyr18B, forming tight aromatic π–π stacks like the pyridine ring of 1. Additionally, the nitrogen atoms (N1) in the azaindole rings of 3a and 3b could interact with Asp219 of NAMPT by a hydrogen bond (Figure 4b), thus suggesting that they might be involved in the tighter specific binding of 3a and 3b to NAMPT than that of 1.

However, this specific contacts of the head-azaindole moiety with NAMPT could not explain the discrepancy in the considerably weaker inhibitory activity of 3b than that of 3a or 1. Therefore, to better understand this phenomenon, we focused on the importance of linker-motifs for the potencies of NAMPT inhibitors. Thus, we analyzed closely the binding forms of the linker-azacyclohexane motifs of 3a and 3b in the tunnel cavity of NAMPT. The carbon atom (C4) of piperidine in 3a and nitrogen atom (N1) of piperazine in 3b were revealed to protrude approximately 1Å deeper into the NMA binding active site, as compared to the corresponding carbon atom of piperidine in 1. Importantly, the electronic repulsion of the replaced nucleophilic N1 atom of piperazine in 3b with the Nδ atom of the imidazole side chain of His191, located 3.82 Å away from the N1 atom, was revealed to occur (Figure 4c). In contrast, there was no electronic repulsion of the corresponding carbon atom of piperidine in 3a with the Nδ atom as well as that of 1. These results clearly show that the electronic repulsion interferes with the proper binding of 3b to the tunnel cavity of NAMPT.

It is noteworthy that Zheng and colleagues have reported that the imidazole ring of His191 interacts with the linker aromatic group of GNE series inhibitors through a herringbone stacking [44,45,47]. Furthermore, NAMPT mutations, including H191R, D93del, Q388R, S165Y and G217V, have been reported to confer resistance to NAMPT inhibitors [50–52]. In addition, we have recently shown that an NAMPT mutation variant of HCT116 cells generated by continuously exposing cells to increasing concentrations of 1, which confers cross-resistance to diverse NAMPT inhibitors, such as CHS-828, GNE-617 and STF-118804, was mapped to only H191R [36,37]. Thus, this His191 residue should be considered with higher priority in designing further effective NAMPT inhibitors. That is, it is preferable to avoid interactions with His191 in designing novel NAMPT inhibitors.
Figure 4. Binding modes of 3a, 3b and 1 in the tunnel cavity of NAMPT. (a) Binding modes of 3a, 3b and 1 in the tunnel cavity of NAMPT were analyzed by our in silico binding mode analysis using RDKit [48] and LigandScout [49]. (b) Close-up view of head regions of these three NAMPT inhibitors in the tunnel cavity of NAMPT using Open3DALIGN [53]. (c) Close-up view of the binding interactions of azacyclohexane-linker moieties of these NAMPT inhibitors with His191A located in the tunnel cavity of NAMPT. Carbons are colored green (3a), magenta (3b) and gray (1). Nitrogen and oxygen are colored blue and red, respectively.
2.3. Influence of Enthalpic Effects of 3a and 3b on the Interactions with Tunnel Cavity of NAMPT

The C-to-N conversion in the azacyclohexane-linker moiety was considered to be possible to introduce some entropic differences in the compounds’ abilities to interact with the tunnel cavity of NAMPT. Although piperazine and piperidine rings are known to form various conformations from chairs to half-chair and skewed boat-to-boat forms in aqueous solution [54], piperazine-containing 3b and piperidine-containing 3a took nearly identical positions as a chair form to that occupied by 1 in the tunnel cavity analyzed by in silico binding modes and crystal structure analyses [38–42] (Figure 4). In addition, both C4 in 3a (Figure 5a) and tertiary amine-type nitrogen (N1) containing 3b (Figure 5b) might prefer equatorial conformations to axial ones by 1,3-diaxial hydrogen interaction [54], taking a stable chair conformation in the tunnel cavity of NAMPT. Therefore, such entropic effects in 3a and 3b in the tunnel space of NAMPT may not be so different from each other. Thus, it is possible to consider that the dramatic differences in enzymatic and cellular potencies between 3a and 3b may be due to enthalpic effects rather than entropic effects inside the tunnel cavity of NAMPT.

To prove this hypothesis, we attempted to analyze the binding affinity scores of 3a and 3b to NAMPT (PDB Code: 2GVJ) by our in silico binding mode analyses using RDKit [48] and LigandScout [49], as described in Materials and Methods. Remarkably, the binding affinity scores of 3a and 3b (non-protonation form) (Figure 5b) were calculated to be $-29.07$ and $-22.13$ kJ/mol, respectively, while that of 1 ($-28.05$ kJ/mol) was near to the value of 3b (Table 1). Thus, the piperidine-motif was revealed to be beneficial for 3a potency and has valuable influences on the interaction with the tunnel cavity of NAMPT.

The N1 and N4 atoms in 3a and 3b, respectively, would take an $sp^2$-like bond, due to the effect of the proximity of the benzoyl group (R2) on the lone pair (Figure 5a,b). In contrast, although it is hard to know whether or not the N1 atom of piperazine in 3b, which takes an $sp^3$ bond like the C4 atom in 3a, would be protonated in the tunnel cavity of NAMPT, it is possible to be protonated in physiological conditions. Thus, the binding affinity score of 3b in the protonation state was measured by LigandScout. Interestingly, it was calculated to be $-23.85$ kJ/mol and was not as good as that of the non-protonated form (Table 1). In the protonated N1 atom, the electrostatic interaction becomes attractive to the Nδ of His 191 but repulsive against the protonated Nδ. However, as the imidazole ring of His has aromaticity, the positive charge introduced by protonation does not tend to be localized on the Nδ. Thus, the repulsion may not be so strong. In addition, it seems that a hydrogen bond between the protonated Nδ of His191 and the N1 atom in 3b could not be formed, because the angle that NδH-N1 (piperazine) takes is 106.4°. Furthermore, of course, the influence of the hydrophobic amino acids in the tunnel region on the protonated N1 in 3b cannot be ruled out, and they may negatively influence this interaction. Accordingly, the electrostatic enthalpy effects were suggested to interfere with the proper binding of 3b to the tunnel cavity of NAMPT. For this reason, both the non-protonated and protonated forms of 3b are suggested to have lower binding affinities to the tunnel cavity of NAMPT than does 3a. These outcomes indicate that the electrostatic potential (enthalpy term) inside the tunnel region of NAMPT has a significant impact on the different binding affinity of 3b from that of 3a.
Figure 5. Stable conformers of 3a and 3b in the tunnel cavity of NAMPT. The conformational states of 3a (a) and 3b with its protonation form (b) in the tunnel cavity of NAMPT were analyzed by in silico binding mode analysis and crystal structure of 1 (PDB code: 2GVJ) [38–42]. The hydrogen interactions of axial-methylene with 1,3-diaxial hydrogens are represented by dashed lines.

Table 1. Biochemical properties and binding affinity scores of 3a and 3b NAMPT inhibitors.

| Compound | IC₅₀ a (μM) | Binding Affinity Score b (kJ/mol) |
|----------|-------------|-----------------------------------|
| 3a       | 0.11        | −29.07                            |
| 3b       | 5.06        | −22.13                            |
| 1        | 0.52        | −28.05                            |

a The IC₅₀ values of these three NAMPT inhibitors were determined by the titration curves in Figure 3a. b The binding affinity scores of these three inhibitors in the tunnel cavity of NAMPT were calculated by LigandScout [49].
3. Materials and Methods

3.1. Enzymatic NAMPT Assay

For the inhibitory effects of 3a and 3b on NAMPT activity, recombinant NAMPT activity was measured using a coupled-enzyme reaction system (CycLex NAMPT Colorimetric Assay Kit: MEDICAL & BIOLOGICAL LABORATORIES CO., LTD., Nagoya, Japan) according to the manufacturer’s instructions, as described previously [31,36,37]. Briefly, NAMPT enzyme and tested compounds were put on the 96-well transparent plate. The mixture of the rest of contents (ATP, NAM, PRPP, nicotinamide nucleotide adenylyltransferase 1 (NMNAT1), water-soluble tetrazolium salts (WST-1), alcohol dehydrogenase, diaphorase and ethanol (EtOH)) were then added onto the wells. After a brief incubation, the absorbance of the samples was detected at 450 nm on every 10 min until 60 min using a SYNERGY HTX (BioTek Instruments, Inc.; Winooski, VT, USA). The inhibition rates were calculated by the initial slope of the absorbance of the tested samples divided by that of the control wells. The IC\textsubscript{50} values for the samples were calculated by 4-parameter logistic regression using Image J 1.48v (https://imagej.nih.gov/ij/).

3.2. Cells and Cell Culture

HCT116 cell was obtained from American Type Culture Collection (ATCC) (Manassas, US) and cultured in Dulbecco’s modified Eagle medium (DMEM) (FUJIFILM Wako Pure Chemical Corporation) supplemented with 10% of fetal bovine serum (Biosera Europe; Nuaillé, France), 100 Units/mL of penicillin G and 100 µg/mL of Streptomycin (FUJIFILM Wako Pure Chemical Corporation) [36,37].

3.3. Cell Activity Assay

One thousand HCT116 cells were inoculated into the 96-well transparent cell culture plate and cultured for 24 h. The medium was changed to new medium that contained the tested compounds and was further cultivated for 72 h. Following the culture, 1/10 volume of the WST-8 reagent (FUJIFILM Wako Pure Chemical Corporation) was added and incubated for 1 h. After the incubation, the absorbance of the samples was detected at 450 nm using a SYNERGY HTX (BioTek Instruments, Inc.). Cell activity was expressed as percentage of control cells treated with vehicle alone [31,36,37].

3.4. In Silico Binding Mode and Binding Affinity Score Analyses

The crystal structure of NAMPT was downloaded from the Protein Data Bank (PDB code: 2GVJ) for molecular binding mode analyses. In silico binding mode analyses were done using RDKit [48] and LigandScout [49]. First, 1000 conformations of each inhibitor were generated using the EmbedMultipleConfs and MMFFGetMoleculeForceField functions of RDKit. Second, the conformations of each inhibitor were aligned to the crystal structure of NAMPT using Open3DALIGN [53] implemented in RDKit. The conformation of each inhibitor, which has the highest alignment score, was selected. Third, energy minimization of the selected conformation of each inhibitor with NAMPT was performed, and then the binding affinity score of each inhibitor was calculated using LigandScout [49].

3.5. Synthetic Procedures of the Target Compounds

3.5.1. General Experimental Methods

\textsuperscript{1}H-NMR spectra were recorded at 400 MHz using a JEOL ECZ400 operating (JEOL Ltd.; Tokyo, Japan) at the indicated frequencies. Chemical shifts were expressed in ppm relative to the internal standard tetramethylsilane (ppm = 0.00). All reagents and solvents were purchased from FUJIFILM Wako Pure Chemical Corporation. The progress of all reactions was monitored by TLC using ethyl acetate/hexane, acetone/hexane, dichloromethane (CH\textsubscript{2}Cl\textsubscript{2})/MeOH and CH\textsubscript{2}Cl\textsubscript{2}/MeOH/triethylamine (TEA) as the solvent system, and spots were visualized by irradiation with ultraviolet light (254 nm) or ninhydrin reaction. Column chromatography was performed using silica gel (200–300 mesh).
3.5.2. General Procedures for Synthesizing the Target Compounds 3a and 3b

Compounds 1-benzoylpiperidin-4-yl)butyl)isoindoline-1,3-dione (1a) and 2-(4-(4-benzoylpiperazin-1-yl)butyl)isoindoline-1,3-dione (1b) were synthesized according to Gilig et al. [55] with slight modifications.

3.5.3. (4-(4-aminobutyl)piperidin-1-yl)(phenyl)methanone (2a)

Hydrazine hydrate (0.3 mL, 6.17 mmol) was added under Ar atmosphere to a solution of 1a (1.0 g, 2.56 mmol) in EtOH at 20 °C. After stirring at 20 °C for 10 min, the mixture was heated under reflux for 2 h. After cooling to 20 °C, the white precipitate was filtered off and washed with EtOH (10 mL). The EtOH solutions were combined, and the solvent was evaporated in vacuo. The residue was taken in CH2Cl2 (20 mL) and a saturated aqueous solution of K2CO3 (20 mL). Vigorous stirring for 10 min produced two clear phases. The aqueous phase was extracted with CH2Cl2 (10 mL, 3 times). The combined organic phases were washed with brine (30 mL) and dried over Na2SO4. After solvent evaporation in vacuo, 2a was obtained (0.530 g) as a pale-yellow oil.

3.5.4. N-(4-(1-benzoylpiperidin-4-yl)butyl)-1H-pyrrolo[3,2-c]pyridine-2-carboxamide (3a)

First, 1H-pyrro[3,2-c]pyridine-2-carboxylic acid (0.149 g, 0.918 mmol) was added to a solution of 2a obtained above (0.239 g, 0.918 mmol) in dry DMF (8.0 mL) in a 25 mL round-bottomed flask. HOBt (0.186 g, 1.38 mmol), EDC-HCl (0.352 g, 1.84 mmol) and N,N-diisopropylethylamine (0.462 mL, 2.75 mmol) were added sequentially, and the suspension was allowed to stir at room temperature overnight. The reaction mixture was then poured into a separatory funnel containing water and ethyl acetate. The aqueous phase was separated and extracted three times with EtOAc. The combined organic phase was washed with brine (30 mL) and dried over Na2SO4. After solvent evaporation in vacuo, 3a was obtained (0.215 g, yield 58%, more than 95%, as judged by ODS(C18) HPLC) as white crystals.

3.5.5. 4-(4-Aminobutyl)piperazin-1-yl)(phenyl)methanone (2b)

Compound 2b was obtained from 1b by similar synthetic processes as described in Section 3.5.3.

3.5.6. N-(4-(4-Benzoylpiperazin-1-yl)butyl)-1H-pyrrolo[3,2-c]pyridine-2-carboxamide (3b)

The procedure was the same as described above for the synthesis of 3a. Compound 3b was obtained as pale-yellow powder (0.104 g, yield 28%, more than 95%, as judged by ODS(C18) HPLC). 1H-NMR (DMSO-d6) δ 11.97 (s, broad, 1H), 8.90 (s, 1H), 8.61 (t, 1H), 8.19 (d, 1H), 7.39 (m, 3H), 7.31 (m, 3H), 7.20 (s, 1H), 3.26–3.58 (m, 9H), 2.30–2.38 (m, 5H), 1.48–1.55 (m, 3H).

3.6. Statistical Analysis

All data were obtained from at least three independent experiments and were expressed as mean ± standard error (SE).

4. Conclusions

In this study, to understand the structural basis of beneficial design for effective NAMPT inhibitors, we designed and synthesized a close structural analogue set of potent and specific azacyclohexane-motif NAMPT inhibitors, namely 3a (piperidine-motif) and 3b (piperazine-motif) modified from 1. In addition, these two compounds have 5-azaindole ring as the head structure, which is replaced by the vinylpyridine ring of 1. Through biochemical experiments and in silico binding mode analyses using these inhibitors, we clearly showed that 3a is the most potent NAMPT inhibitor among these inhibitors, and that the azaindole-head and piperidine-linker of NAMPT inhibitors are promising
motifs that result in enthalpically higher potent inhibitory activity against NAMPT molecule and cancer cells. The main reason for this phenomenon was revealed to be due to the electronic repulsion of the piperazine-linker motif of 3b with His191 of NAMPT. Importantly, this H191 is known to be a drug-resistance sensitive amino acid residue [36,37,50–52]. Thus, it is better to avoid interactions with His191 in designing effective NAMPT inhibitors. Further studies on the tunnel cavity of NAMPT using rigid analogues of azaindole-piperidine-motif inhibitors will allow a more definitive understanding of the inhibitor pharmacophore. Although additional confirmatory studies, such as the synthesis of novel azaindole-piperidine-motif NAMPT inhibitors and details of in vivo efficacy and resistibility are warranted, our findings provide valuable insights into the design for effective NAMPT inhibitors that offer improved therapeutic potential by making high specificity and avoiding resistance to NAMPT. Furthermore, on the basis of the characteristics of NAMPT functions and the mechanisms of action of NAMPT inhibitors, we are now trying to design additional azaindole–piperidine-motif NAMPT inhibitors available in vivo, such as the translation of a potent one into a payload for antibody–drug conjugates, which are an important therapeutic modality enabling targeted drug delivery to cancer cells [56].

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