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Structural Modifications of DAPY Analogues with Potent Anti-HIV-1 Activity

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A novel series of diarylpyrimidine analogues (DAPYs) featuring a naphthyl moiety at the C4 position were synthesized and evaluated for their in vitro activity against HIV in MT-4 cells. All compounds exhibited strong activity against wild-type HIV-1. The most active compound showed activity against wild-type HIV-1

with an EC₅₀ value of 2.35 nm and against the double mutant strain (K103N+Y181C) with an EC₅₀ value of 6.6 μ m, with a selectivity index greater than 60000 against wild-type HIV-1. Additionally, some compounds also showed activity against HIV-2 (EC₅₀= 5.82 μ m).

Introduction

Since they were first reported in 2001,^[1] diarylpyrimidine derivatives (DAPYs), a novel class of non-nucleoside reverse transcriptase inhibitors (NNRTIs), have attracted considerable attention due to their excellent potency against wild-type and mutant strains of HIV-1 reverse transcriptase (HIV-1 RT), relative to other NNRTIs.^[2-6] Further modifications of DAPYs have focused on wing I substituents and the pyrimidine scaffold, leading to the synthesis of many promising lead compounds^[7-9] with strong potency against drug-resistant virus strains. Among these, etravirine (TMC125, **2**, Figure 1) has been approved by the U.S. Food and Drug Administration (FDA) for the treatment of patients infected with HIV-1 variants that are resistant to other anti-retroviral drugs.^[10]

Our long-standing work on NNRTI pyrimidine analogues such as HEPTs (3),^[11] DABOs (4),^[12] DATAs (5)^[13,14] and DAPYs^[15] (Figure 1) has indicated that replacement of the phenyl ring at the C6 position of the pyrimidine ring with a bulky naphthyl moiety is beneficial by improving the π – π stacking interactions between inhibitors and amino acid residues Tyr 181, Tyr 188,

Figure 1. Chemical structures of DAPYs, HEPTs, DABOs, DATAs, and etravirine.

and Trp 229 within the binding pocket of RT. Herein we report the synthesis, antiviral activity, and preliminary structure–activity relationships (SARs) of these new naphthyl-substituted DAPYs.

Results and Discussion

Chemistry

Target compounds **10 a–af** in this study were synthesized as depicted in Scheme 1. Key intermediate 2-(methylthio)pyrimidin-4(1*H*)-ones **7 a–c** were readily prepared by S-alkylation of thiouracils **6 a–c** with iodomethane according to our previous reported protocol. Compounds **7 a–c** were condensed with 4-cyanoaniline at 180–190 °C for 8 hours under solvent-free conditions to afford 4-(4-oxo-1,4-dihydropyrimidin-2-ylamino)-benzonitriles **8 a–c.** Subsequent treatment with POCl₃ at reflux for 30 minutes provided the corresponding 4-(4-chloropyrimidin-2-ylamino)benzonitriles **9 a–c.** Treatment of **9 a–c** with appropriate naphthol or naphthiol derivatives in the presence of K₂CO₃ in anhydrous DMF at 110 °C under nitrogen atmosphere gave desired compounds **10 a–af** in yields ranging from 20.1 to 87.9%.

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Scheme 1. Synthetic route to compounds 10 a-af. Reagents and conditions: a) NaOH, room temperature, 30 min, then Mel, room temperature, 24 h; b) 4-cyanoaniline, $180-190^{\circ}$ C, ~ 8 h; c) POCl₃, reflux, 30 min; d) potassium carbonate, naphthol or naphthol derivative, DMF, 5 min, then 9, 110° C, N_2 , 8-12 h.

Biological activity

The MTT method^[17,18] was used to evaluate 29 new naphthyl-substituted compounds, **10 a–10 ac**, and 3-phenylsubstituted compounds, **10 ad–10 af**, along with three FDA-approved drugs for reference purposes: nevirapine (NEV), delavirdine (DEV), and efavirenz (EFV). These compounds were all assessed for their cytotoxicity and antiviral activities in MT-4 cells infected with the HIV-1 wild-type virus (LAI strain, IIIB), HIV-1 double mutant virus (K103N+Y181C, with Lys103 replaced by Asn and Tyr181 by Cys), or HIV-2 strain ROD.

Two series of DAPY derivatives: 1-naphthyl-substituted compounds 10 a-j and 2-naphthyl-substituted compounds 10 k-ac, were generally very active against wild-type HIV-1 and did not exhibit cytotoxicity up to 24.86 µm, with the exception of 10 b, as shown in Table 1. The anti-retroviral activity (EC₅₀) ranged from 2.35 nm (10 w) to 111.53 nm (10 aa); the selectivity index (SI) ranged from 309 (10 h) to > 106232 (10 t). The 1-naphthyland 2-naphthyl-substituted parent compounds 10c-d, 10m, and 10p were more potent than NEV and DEV, and demonstrated inhibitory potency in the low-nanomolar range with high SI values. With the aim of validating our design rationale, three compounds, 10 ad-af, incorporating a phenyl ring at the C4 position of the pyrimidine ring were synthesized. These compounds exhibited poor activity against the HIV-1 LAI virus, and were 3- to 12-fold less potent than the naphthyl-substituted compounds 10 c-d, 10 m, and 10 p. These results confirmed our initial hypothesis that replacement of the phenyl moiety with a naphthyl moiety improves the putative π – π stacking interactions between inhibitors and amino acid residues Tyr 181, Tyr 188, and Trp 229 within the binding pocket of RT.

To investigate structural variations of the pyrimidine and naphthalene rings, a series of DAPYs with substitutions at the C4 position of these rings was synthesized and evaluated for antiviral activity. Introduction of a methyl group on the pyrimidine ring for the 1-naphthyl or 2-naphthyl series resulted in negligible loss of anti-HIV-1 (wild-type) activity. On the other hand, it is interesting to note that 10b, 10f, and 10i, with methyl substituents at the C5 position, were more potent than the corresponding nonsubstituted compounds in the 1-naphthyl series. However, in the 2-napthyl series, 10 m, 10 w, and 10 z, which lack substituents at the C5 position, exhibited the highest potency. Introduction of a chlorine or bromine at the para position (10e-j) resulted in minor changes in activity as compared with 10a-c for the 1-naphthyl series. Conversely, in the 2-naphthyl series, the 1-chloro- or bromo-substituted compounds 10 r-w showed significant improvements in potency compared with original hits 10k-m, with 10w displaying the greatest activity (EC₅₀ = 2.35 nm).

To investigate the potency of these compounds against drug-resistant virus, the activities against the double mutant strain K103N+Y181C were also evaluated. The compounds with chlorine or bromine substituents at the C1 position in the 2-naphthyl series exhibited low-micromolar antiviral activity

Compd	EC ₅₀ [nм] ^[а] wild-type (IIIB)	ЕС ₅₀ [μм] ^[а] HIV-2	EC ₅₀ [μм] ^(a) K103N + Y181C	CC ₅₀ [µм] ^[b]	SI ^[c]
10 a	68.96 ± 36.04	> 34.56	> 34.56	34.56 ± 1.79	501
10 b	3.35 ± 0.45	> 4.06	> 4.06	4.06 ± 1.53	1220
10 c	21.87 ± 13.39	> 369.43	> 369.43	> 369.43	> 16 903
10 d	$\textbf{5.87} \pm \textbf{3.72}$	16.17	> 252.18	252.18 ± 24.55	43 072
10 e	60.49 ± 12.41	> 58.22	> 58.22	58.22 ± 18.04	964
10 f	$\textbf{9.33} \pm \textbf{1.78}$	> 323.14	> 323.14	> 323.14	> 34 626
10 g	12.15 ± 0.08	> 335.29	> 335.29	> 335.29	> 27 594
10 h	80.46 ± 30.84	> 24.86	> 24.86	24.86 ± 1.18	309
10 i	10.39 ± 1.07	> 212.53	> 212.53	212.53 ± 26.32	20 445
10 j	23.97 ± 6.95	> 203.71	> 203.71	> 203.71	≥8492
10 k	54.77 ± 1.14	> 14.02	> 156.3	156.3 ± 61.32	2861
101	9.96 ± 1.48	>354.72	> 354.72	> 354.72	> 35 663
10 m	9.78 ± 2.19	223.14	> 369.43	> 369.43	> 37 764
10 n	47.22 ± 9.50	7.06	> 30.32	30.32 ± 5.75	642
10 o	24.4 ± 15.14	> 251.87	> 251.87	> 251.87	> 10 323
10 p	12.70 ± 1.41	≥ 10.13	>47.20	47.20 ± 2.82	3754
10 q	49.16 ± 51.23	18.35	> 129.77	129.77 ± 82.59	2636
10 r	11.40 ± 0.75	6.08	4.27	77.35 ± 53.46	6790
10 s	3.83 ± 1.22	> 323.14	7.55	> 323.14	>84650
10 t	3.17 ± 0.40	168.77	52.84	> 335.29	> 106 232
10 u	8.42 ± 2.67	5.82	4.52	38.95 ± 7.23	4634
10 v	4.57 ± 0.60	6.72	9.51	> 289.83	>63613
10 w	2.35 ± 0.07	15.94	6.57	153.5 ± 10.40	65 591
10 x	18.42 ± 13.33	≥ 50.18	≥ 213.65	≥ 213.65	≥ 11 602
10 y	12.96 ± 6.47	75.07	> 245.01	_ > 245.01	> 18 925
10 z	11.67 ± 3.06	136.15	> 251.94	> 251.94	> 21 608
10 aa	111.53 ± 57.50	> 230.82	> 230.82	230.82 ± 42.78	2070
10 ab	85.56 ± 32.00	> 289.83	> 289.83	> 289.83	> 3391
10 ac	25.4 ± 22.29	> 299.57	> 299.57	> 299.57	> 11 767
10 ad	262.3 ± 202.10	> 21.00	> 21.00	21 ± 14.12	80
10 ae	37.71 ± 20.18	> 0.30	> 0.30	$\textbf{0.3} \pm \textbf{0.05}$	8
10 af	70.76 ± 28.79	> 433.58	> 433.58	> 433.58	>6137
NEV	75.1	_	_	> 15.02	> 252
EFV	3	_	_	> 6.336	> 2174
DEV	72	_	_	> 3.619	44

[a] EC_{50} : compound concentration required to protect the cell against viral cytopathogenicity by 50% in MT-4 cells. [b] EC_{50} : compound concentration that decreases the normal uninfected MT-4 cell viability by 50%. [c] SI: selectivity index; ratio EC_{50} / EC_{50} (wild-type).

against the double mutant strain: ($10 \, s$,EC₅₀= $7.55 \, \mu m$; $10 \, r$, EC₅₀= $4.27 \, \mu m$; $10 \, u$, EC₅₀= $4.52 \, \mu m$; $10 \, v$, EC₅₀= $9.51 \, \mu m$; and $10 \, w$, EC₅₀= $6.57 \, \mu m$). These promising results promote further investigation of this new series of NNRTIs.

All title compounds were also assessed for their abilities to inhibit replication of the HIV-2 ROD virus in MT-4 cells, as depicted in Table 1. Some compounds also displayed activity against HIV-2 at micromolar concentrations, particularly $10\,n$ (EC $_{50}=7.06\,\mu\text{M}$), $10\,r$ (EC $_{50}=6.08\,\mu\text{M}$), $10\,u$ (EC $_{50}=5.82\,\mu\text{M}$), $10\,v$ (EC $_{50}=6.72\,\mu\text{M}$), and $10\,w$ (EC $_{50}=15.94\,\mu\text{M}$); however, these compounds were also fairly cytotoxic to the host cells.

Molecular modeling analysis

To investigate the possible binding conformations of our newly synthesized compounds and their interaction mode with RT, a modeling study was performed using the AutoDock 4.0.1 program. Compound **10 m** was chosen to be docked into the non-nucleoside inhibitor binding pocket (NNIBP) of HIV-1 RT. Coordinates of the NNIBP were taken from a crystal structure of the RT-TMC278 complex, owing to the high

degree of similarity between TMC278 and $10\,m$. ^[20] The theoretical binding mode of $10\,m$ to the NNIBP is shown in Figure 2. In comparison with the binding of TMC125 to RT, ^[21] the naphthyl-substituted DAPY $10\,m$ enhanced π - π and π -H interactions with amino acid residues Tyr181, Tyr188, and Phe 227, over those of the original cyanovinyl group in TMC278. ^[20] These binding data support our initial design; however, these compounds lack the strong interactions with Trp 229 that play an important role in the activity of TMC278 and TMC125 against drug-resistant mutant strains.

Conclusions

In summary, we designed and synthesized a novel series of naphthyl-substituted DAPYs. Biological test results indicated that the designed compounds showed potent antiviral activity, with EC $_{50}$ values in the low-nanomolar range, and did not exhibit cytotoxicity up to 24860 nm, with the exception of 10 b. The 1-bromo-2-naphthoxy compound 10 w was the most potent inhibitor of wild-type HIV-1 (EC $_{50}$ =2.35 nm in MT-4 cells); however, the activity of 10 w against the double mutant

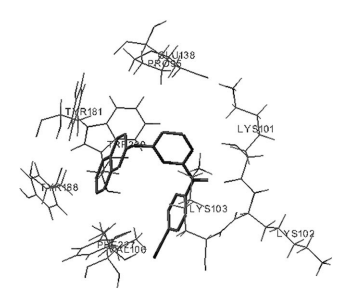


Figure 2. Model of $10\,\mathrm{m}$ docked within the non-nucleoside binding site of RT.

strain K103N+Y181C were still at the micromolar level. These results serve to support further modification of the naphthalene ring in an attempt to improve activity against drug-resistant mutant strains.

Experimental Section

Chemistry. Melting points were measured on a WRS-1 digital melting point apparatus and are uncorrected. ^1H NMR spectra were recorded in [D₆]DMSO using a Bruker AV 400 MHz spectrometer. Chemical shifts are reported in δ (ppm) units relative to the internal standard tetramethylsilane (TMS). Mass spectra were obtained on an Agilent MS/5975 mass spectrometer. Elemental analyses were performed on a Carlo Erba 1106 instrument, and the results of elemental analyses for C, H, Cl, N, and S were within $\pm 0.4\%$ of theoretical values. All chemicals and solvents used were of reagent grade and were purified and dried by standard methods before use. All air-sensitive reactions were run under a nitrogen atmosphere. All reactions were monitored by TLC on pre-coated silica gel G plates at 254 nm under a UV lamp using EtOAc/hexanes as eluents. Flash chromatography separations were obtained using silica gel (300–400 mesh).

General procedure for the synthesis of $7\,a$ –c. NaOH (8.0 g, 300 mmol) was added portion-wise to a suspension of thiouracils $6\,a$ –c (300 mmol) in H_2O at room temperature. After the reaction mixture was stirred for 30 min, iodomethane (320 mmol) was added, and stirring continued at room temperature for 24 h. The precipitate was filtered off, washed with H_2O , and dried to give 2-(methylthio)pyrimidin-4(1H)-ones $6\,a$ –c to be used without further purification.

General procedure for the synthesis of $8\,a$ –c. 2-(methylthio)pyrimidin-4(1H)-ones $7\,a$ –c (200 mmol) and 4-cyanoaniline (500 mmol) were thoroughly mixed. The mixture was slowly heated to 180– $190\,^{\circ}C$ and maintained at this temperature until no odor of methanethiol was perceptible (about $8\,h$). After cooling and dissolving the mixture in EtOH, and subsequent decolorization with charcoal, the product was precipitated with H_2O . The suspension was acidified (pH 3) with HCI to dissolve remaining starting materials. The

precipitate was filtered and washed with H_2O . Final products were crystallized from 90% EtOH.

General procedure for the synthesis of 9 a–c. A mixture of 40 mL POCl₃ and intermediate 4-(4-oxo-1,4-dihydropyrimidin-2-ylamino)-benzonitriles **8 a–c** (150 mmol) was held at reflux for 30 min. The mixture was poured into 250 mL ice-water and stirred at room temperature for 3 h. The resulting precipitate was filtered, washed with 50 mL H₂O, and dried to give 4-(4-chloropyrimidin-2-ylamino)-benzonitriles **9 a–c** to be used without further purification.

General procedure for the synthesis of 10 a–af. Potassium carbonate (10 mmol) was added to a solution of naphthol or naphthiol derivatives (2 mmol) in 20 mL anhydrous DMF and stirred for 5 min. One of the 4-(4-chloropyrimidin-2-ylamino)benzonitriles 9 a–c (2 mmol) was then added. The mixture was heated at 110 °C under nitrogen atmosphere for 8–12 h. Next, the mixture was treated with cold $\rm H_2O$ (200 mL), and the resulting precipitate was collected by filtration. Crude products 10 a–af were recrystallized from toluene.

4-(4-methyl-6-(naphthalen-1-yloxy)pyrimidin-2-ylamino)benzonitrile (10 a). Yield 23.2 %; recrystallized from toluene, mp: 142.8–143.8 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.39 (s, 3 H, CH₃), 6.62 (s, 1 H, CH), 7.31–7.41 (m, 4 H, Ph), 7.43–8.10 (m, 7 H, Naph), 10.06 ppm (s, 1 H, NH); MS (EI) m/z: 352 [M] +; Anal. (C₂₂H₁₆N₄O) C, H. N.

4-(5-methyl-4-(naphthalen-1-yloxy)pyrimidin-2-ylamino)benzonitrile (10 b). Yield 23.9 %; recrystallized from toluene, mp: 178.6–181.5 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.37 (s, 3 H, CH₃), 7.24–7.29 (m, 4 H, Ph), 7.43–8.10 (m, 7 H, Naph), 8.38 (s, 1 H, CH), 9.86 ppm (s, 1 H, NH); MS (EI) m/z: 352 [M] +; Anal. (C₂₂H₁₆N₄O) C, H, N.

4-(4-(naphthalen-1-yloxy)pyrimidin-2-ylamino)benzonitrile (10 c). Yield 22.1%; recrystallized from toluene, mp: 205.7–207.4 °C;

¹H NMR (DMSO, 400 MHz) δ = 6.75 (d, 1 H, J = 6.0 Hz, CH), 7.34–7.42 (m, 4 H, Ph), 7.44–8.01 (m, 7 H, Naph), 8.49 (d, 1 H, J = 6.0 Hz, CH), 10.09 ppm (s, 1 H, NH); MS (EI) m/z: 338 $[M]^+$; Anal. (C₂₁H₁₄N₄O) C, H, N.

4-(4-(naphthalen-1-ylthio)pyrimidin-2-ylamino)benzonitrile

(10 d). Yield 35.2%; recrystallized from toluene, mp: 208.0–208.6 °C; ^1H NMR ([D_6]DMSO, 400 MHz) $\delta\!=\!6.52$ (d, 1 H, $J\!=\!6.0$ Hz, CH), 7.34 (s, 4 H, Ph), 7.56–8.28 (m, 7 H, Naph), 8.19 (d, 1 H, $J\!=\!4.4$ Hz, CH), 10.16 ppm (s, 1 H, NH); MS (EI) m/z: 354 $[M]^+$; Anal. (C_21H14N4S) C, H, N, S.

4-(4-(4-chloronaphthalen-1-yloxy)-6-methylpyrimidin-2-ylamino)-benzonitrile (10 e). Yield 27.3 %; recrystallized from toluene, mp: 195.7–197.0 °C; ^1H NMR ([D₆]DMSO, 400 MHz) $\delta = 2.39$ (s, 3 H, CH₃), 6.65 (s, 1 H, CH), 7.33–7.43 (m, 4 H, Ph), 7.45–8.28 (m, 6 H, Naph), 10.06 ppm (s, 1 H, NH); MS (EI) m/z: 386 [*M*] $^+$; Anal. (C₂₂H₁₅N₄OCl) C, H, N, Cl.

4-(4-(4-chloronaphthalen-1-yloxy)-5-methylpyrimidin-2-ylamino)-benzonitrile (10 f). Yield 31.0%; recrystallized from toluene, mp: 247.2–247.9 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.36 (s, 3 H, CH₃), 7.26–7.33 (m, 4 H, Ph), 7.46–8.30 (m, 6 H, Naph), 8.39 (s, 1 H, CH), 9.87 ppm (s, 1 H, NH); MS (EI) m/z: 386 $[M]^+$; Anal. (C₂₂H₁₅N₄OCI) C, H, N, Cl.

4-(4-(4-chloronaphthalen-1-yloxy)pyrimidin-2-ylamino)benzonitrile (10 g). Yield 47.1%; recrystallized from toluene, mp: 220.8–221.6 °C; ¹H NMR (DMSO, 400 MHz) δ =6.79 (d, 1 H, J=6.4 Hz, CH), 7.37–7.45 (m, 4 H, Ph), 7.47–8.29 (m, 6 H, Naph), 8.50 (d, 1 H, J=

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6.4 Hz, CH), 10.08 ppm (s, 1 H, NH); MS (EI) m/z: 373 $[M]^+$; Anal. $(C_{21}H_{13}N_4OCI)$ C, H, N, CI.

- **4-(4-(4-bromonaphthalen-1-yloxy)-6-methylpyrimidin-2-ylamino)benzonitrile (10 h).** Yield 20.6%; recrystallized from toluene, mp: 193.4–195.3 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.41 (s, 3 H, CH₃), 6.67 (s, 1 H, CH), 7.33–7.41 (m, 4 H, Ph), 7.62–8.25 (m, 6 H, Naph), 10.05 ppm (s, 1 H, NH); MS (EI) m/z: 431[M] +; Anal. (C₂₂H₁₅N₄OBr) C, H, N, Br.
- 4-(4-(4-bromonaphthalen-1-yloxy)-5-methylpyrimidin-2-ylamino)benzonitrile (10 i). Yield 25.3 %; recrystallized from toluene, mp 248.9–250.7 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.34 (s, 3 H, CH₃), 7.26–7.30 (m, 4 H, Ph), 7.38–8.24 (m, 6 H, Naph), 8.37 (s, 1 H, CH), 9.84 ppm (s, 1 H, NH); MS (EI) m/z 431 [M] +; Anal. (C₂₂H₁₅N₄OBr) C, H. N, Br.
- **4-(4-(4-bromonaphthalen-1-yloxy)pyrimidin-2-ylamino)benzonitrile (10 j).** Yield 87.9%; recrystallized from toluene mp: 208.9–209.8 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 6.79 (d, 1 H, J = 6.4 Hz, CH), 7.36–7.44 (m, 4 H, Ph), 7.62–8.25 (m, 6 H, Naph), 8.50 (d, 1 H, J = 6.4 Hz, CH), 10.06 ppm (s, 1 H, NH); MS (EI) m/z: 417 [M]⁺; Anal. (C₂₁H₁₃N₄OBr) C, H, N, Br.
- **4-(4-methyl-6-(naphthalen-2-yloxy)pyrimidin-2-ylamino)benzonitrile (10 k).** Yield 44.4%; recrystallized from toluene, mp: 168.8–170.8 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.38 (s, 3 H, CH3), 6.52 (s, 1 H, CH), 7.25–7.64 (m, 4 H, Ph), 7.41–8.08 (m, 7 H, Naph), 10.10 ppm (s, 1 H, NH); MS (EI) m/z: 352 [M] $^+$; Anal. (C $_{22}$ H $_{16}$ N $_4$ O) C, H, N.
- **4-(5-methyl-4-(naphthalen-2-yloxy)pyrimidin-2-ylamino)benzonitrile (10 I).** Yield 21.9%; recrystallized from toluene, mp: 209.9–211.3 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.26 (s, 3 H, CH3), 7.15–7.53 (m, 4 H, Ph), 7.42–8.08 (m, 7 H, Naph), 8.34 (s, 1 H, CH), 9.91 ppm (s, 1 H, NH); MS (EI) m/z 352 [M] $^+$; Anal. ($C_{22}H_{16}N_4O$) C, H, N.
- 4-(4-(naphthalen-2-yloxy)pyrimidin-2-ylamino)benzonitrile
- **(10 m).** Yield 30.5%; recrystallized from toluene, mp: 204.7–206.4°C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 6.70 (d, 1 H, J = 6.0 Hz, CH), 7.31–7.68 (m, 4 H, Ph), 7.47–8.12 (m, 7 H, Naph), 8.50 (d, 1 H, J = 6.0 Hz, CH), 10.16 ppm (s, 1 H, NH); MS (EI) m/z: 338 [M] $^+$; Anal. (C₂₁H₁₄N₄O) C, H, N.
- **4-(4-methyl-6-(naphthalen-2-ylthio)pyrimidin-2-ylamino)benzonitrile (10 n).** Yield 41.8%; recrystallized from toluene, mp: 180.7–181.6 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.37 (s, 3 H, CH3), 6.66 (s, 1 H, CH), 6.94–7.41 (m, 4 H, Ph), 7.63–8.31 (m, 7 H, Naph), 10.11 ppm (s, 1 H, NH); MS (EI) m/z: 368 $[M]^+$; Anal. (C₂₂H₁₆N₄S) C, H, N, S.
- **4-(5-methyl-4-(naphthalen-2-ylthio)pyrimidin-2-ylamino)benzonitrile (10 o).** Yield 21.4%; recrystallized from toluene, mp: 214.3–215.7 °C; 1 H NMR ([D₆]DMSO, 400 MHz) $\delta\!=\!2.27$ (s, 3 H, CH₃), 6.47–7.05 (m, 4 H, Ph), 7.60–8.18 (m, 7 H, Naph), 8.30 (s, 1 H, CH), 9.91 ppm (s, 1 H, NH); MS (EI) m/z: 368 [*M*] $^{+}$; Anal. (C₂₂H₁₆N₄S) C, H, N, S.
- **4-(4-(naphthalen-2-ylthio)pyrimidin-2-ylamino)benzonitrile** (**10 p).** Yield 27.5 %; recrystallized from toluene, mp: 183.0–183.3 °C;

 ¹H NMR ([D₆]DMSO, 400 MHz) δ = 6.70 (d,, 1 H, J = 5.6 Hz, CH), 7.00–7.45 (m, 4 H, Ph), 7.62–8.26 (m, 7 H, Naph), 8.33 (s, 1 H, CH), 10.18 ppm (s, 1 H, NH); MS (EI) m/z: 354 [M]⁺; Anal. (C₂₁H₁₄N₄S) C, H, N, S.
- **4-(5-bromo-4-(naphthalen-2-ylthio)pyrimidin-2-ylamino)benzonitrile (10 q).** Yield 32.1%; recrystallized from toluene, mp: 193.2–

- 194.1 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 6.46–6.98 (m, 4H, Ph), 7.64–8.35 (m, 7H, Naph), 8.48 (s, 1H, CH), 10.20 ppm (s, 1H, NH); MS (EI) m/z: 433 [M] $^+$; Anal. (C $_{21}$ H $_{13}$ N $_4$ SBr) C, H, N, Br.
- **4-(4-(1-chloronaphthalen-2-yloxy)-6-methylpyrimidin-2-ylamino)-benzonitrile (10 r).** Yield 20.7%; recrystallized from toluene, mp 222.3–224 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ =2.42 (s, 3 H, CH₃), 6.67 (s, 1 H, CH), 7.11–7.48 (m, 4 H, Ph), 7.58–8.23 (m, 6 H, Naph), 10.10 ppm (s, 1 H, NH); MS (EI) m/z: 386 [M] $^+$; Anal. (C₂₂H₁₅N₄OCI) C, H, N, Cl.
- 4-(4-(1-chloronaphthalen-2-yloxy)-5-methylpyrimidin-2-ylamino)-benzonitrile (10 s). Yield 39.3 %; recrystallized from EtOAc, mp: 225.7–227.4 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.30 (s, 3 H, CH₃), 7.06–7.43 (m, 4 H, Ph), 7.59–8.21 (m, 6 H, Naph), 8.37 (s, 1 H, CH), 9.94 ppm (s, 1 H, NH); MS (EI) m/z: 406 [M] †; Anal. (C₂₂H₁₅N₄OCI) C, H, N, CI.
- **4-(4-(1-chloronaphthalen-2-yloxy)pyrimidin-2-ylamino)benzonitrile (10t).** Yield 20.1%; recrystallized from toluene, mp: 201.9–203.2 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 6.79 (d, 1 H, J=5.6 Hz, CH), 7.15–7.51 (m, 4 H, Ph), 7.60–8.23 (m, 6 H, Naph), 8.52 (d, 1 H, J=5.6 Hz, CH), 10.15 ppm (s, 1 H, NH); MS (EI) m/z: 372 [M]⁺; Anal. (C₂₁H₁₃N₄OCI) C, H, N, CI.
- **4-(4-(1-bromonaphthalen-2-yloxy)-6-methylpyrimidin-2-ylamino)benzonitrile (10 u).** Yield 44.5%; recrystallized from toluene, mp: 216.4–217.5 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.41 (s, 3 H, CH₃), 6.65 (s, 1 H, CH), 7.12–7.49 (m, 4 H, Ph), 7.55–8.21 (m, 6 H, Naph), 10.11 ppm (s, 1 H, NH); MS (EI) m/z: 431 $[M]^+$; Anal. ($C_{22}H_{15}N_4OBr$) C, H, N, Br.
- 4-(4-(1-bromonaphthalen-2-yloxy)-5-methylpyrimidin-2-ylamino)benzonitrile (10 v). Yield 35.9%; recrystallized from MeOH, mp: 233.7–236.1 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.31 (s, 3 H, CH₃), 7.07–7.42 (m, 4 H, Ph), 7.59–8.21 (m, 6 H, Naph), 8.39 (s, 1 H, CH), 9.94 ppm (s, 1 H, NH); MS (EI) m/z: 431 [M] †; Anal. (C₂₂H₁₅N₄OBr) C, H, N, Br.
- **4-(4-(1-bromonaphthalen-2-yloxy)pyrimidin-2-ylamino)benzonitrile (10 w).** Yield 33.0%; recrystallized from toluene, mp: 187.6–189.9 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 6.78 (d, 1 H, J = 5.6 Hz, CH), 7.17–7.52 (m, 4 H, Ph), 7.58–8.21 (m, 6 H, Naph), 8.52 (d, 1 H, J = 5.6 Hz, CH), 10.13 ppm (s, 1 H, NH); MS (EI) m/z:417 [M] +; Anal. (C₂₁H₁₃N₄OBr) C, H, N, Br.
- 4-(4-(1,6-dibromonaphthalen-2-yloxy)-6-methylpyrimidin-2-ylamino)benzonitrile (10 x). Yield 53.6%; recrystallized from toluene, mp: 243.2–244.2 °C; ¹H NMR (DMSO, 400 MHz) δ = 2.40 (s, 3 H, CH₃), 6.64 (s, 1 H, CH), 7.19–7.50 (m, 4 H, Ph), 7.48–8.44 (m, 5 H, Naph), 10.09 ppm (s, 1 H, NH); MS (EI) m/z: 510 $[M]^+$; Anal. ($C_{22}H_{14}N_4OBr_2$) C, H, N, Br.
- **4-(4-(1,6-dibromonaphthalen-2-yloxy)-5-methylpyrimidin-2-ylamino)benzonitrile (10 y).** Yield 37.6%; recrystallized from toluene, mp: 240.8–242.1 °C; 1 H NMR ([D₆]DMSO, 400 MHz) $\delta\!=\!2.29$ (s, 3 H, CH₃), 7.16–7.44 (m, 4 H, Ph), 7.64–8.38 (m, 5 H, Naph), 8.46 (s, 1 H, CH), 9.94 ppm (s, 1 H, NH); MS (EI) m/z: 510 [*M*] $^+$; Anal. (C₂₂H₁₄N₄OBr₂) C, H, N, Br.
- **4-(4-(1,6-dibromonaphthalen-2-yloxy)pyrimidin-2-ylamino)benzonitrile** (10 z). Yield 29.9%; recrystallized from toluene, mp: 231.4–232.1 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 6.70 (d, 1 H, J = 5.6 Hz, CH), 7.24–7.54 (m, 4 H, Ph), 7.64–8.46 (m, 5 H, Naph), 8.52 (d, 1 H, J = 5.6 Hz, CH), 10.11 ppm (s, 1 H, NH); MS (EI) m/z: 496 [M] †; Anal. (C₂₁H₁₂N₄OBr₂) C, H, N, Br.

- **4-(4-(6-bromonaphthalen-2-yloxy)-6-methylpyrimidin-2-ylamino)benzonitrile (10 aa).** Yield 22.5 %; recrystallized from toluene, mp: 200.8–201.4 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.29 (s, 3 H, CH₃), 6.63 (s, 1 H, CH), 7.31–7.65 (m, 4 H, Ph), 7.48–8.33 (m, 6 H, Naph), 10.09 ppm (s, 1 H, NH); MS (EI) m/z: 431 $[M]^+$; Anal. ($C_{22}H_{15}N_4OBr$) C, H, N, Br.
- **4-(4-(6-bromonaphthalen-2-yloxy)-5-methylpyrimidin-2-ylamino)benzonitrile (10 ab).** Yield 26.1%; recrystallized from toluene, mp 239.3–240.3 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ = 2.25 (s, 3 H, CH₃), 7.23–7.53 (m, 4 H, Ph), 7.55–8.08 (m, 6 H, Naph), 8.35 (s, 1 H, CH), 9.91 ppm (s, 1 H, NH); MS (EI) m/z: 431 [M]⁺; Anal. (C₂₂H₁₅N₄OBr) C, H, N, Br.
- **4-(4-(6-bromonaphthalen-2-yloxy)pyrimidin-2-ylamino)benzonitrile (10 ac).** Yield 23.7%; recrystallized from toluene, mp: 234.2–236.2 °C; ¹H NMR ([D₆]DMSO, 400 MHz) δ =6.67 (d, 1 H, J=5.6 Hz, CH), 7.34–7.66 (m, 4 H, Ph), 7.51–8.34 (m, 6 H, Naph), 8.48 (d, 1 H, J=5.6 Hz, CH), 10.11 ppm (s, 1 H, NH); MS (EI) m/z: 417 [M] $^+$; Anal. (C $_{21}$ H $_{13}$ N $_4$ OBr) C, H, N, Br.
- **4-(4-methyl-6-phenoxypyrimidin-2-ylamino)benzonitrile (10 ad).** Yield 20.4%; recrystallized from toluene, mp: 138.4–141.4 °C; 1 H NMR ([D₆]DMSO, 400 MHz) $\delta\!=\!2.35$ (s, 3 H, CH₃), 6.43 (s, 1 H, CH), 7.22–7.67 (m, 4 H, Ph), 7.35–7.52 (m, 5 H, Ph), 10.09 ppm (s, 1 H, NH); MS (El) m/z: 302 [M] $^+$; Anal. (C₁₈H₁₄N₄O) C, H, N.
- **4-(5-methyl-4-phenoxypyrimidin-2-ylamino)benzonitrile** (10 ae). Yield 20.3 %; recrystallized from toluene, mp: 170.0–171.5 °C; 1 H NMR ([D₆]DMSO, 400 MHz) δ = 2.20 (s, 3 H, CH₃), 7.23–7.58 (m, 4 H, Ph), 7.36–7.53 (m, 5 H, Ph), 8.29 (s, 1 H, CH), 9.94 ppm (s, 1 H, NH); MS (El) m/z: 302 [M] $^+$; Anal. (C₁₈H₁₄N₄O) C, H, N.
- **4-(4-phenoxypyrimidin-2-ylamino)benzonitrile** (**10 af).** Yield 28.7%; recrystallized from toluene, mp: 188.6–189.7 °C; 1 H NMR ([D₆]DMSO, 400 MHz) δ = 6.67 (d, 1 H, J = 5.6 Hz, CH), 7.25–7.69 (m, 4 H, Ph), 7.35–7.53(m, 5 H, Ph), 8.43 (d, 1 H, J = 5.6 Hz, CH), 10.13 ppm (s, 1 H, NH); MS (EI) m/z: 288 [M] $^{+}$; Anal. (C₁₇H₁₂N₄O) C, H. N.

Evaluation method. Anti-HIV activity and cytotoxicity were evaluated against wild-type HIV-1 strain IIIB in MT-4 cells using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) method. [17,18] Briefly, virus stocks were titrated in MT-4 cells and expressed as the 50% cell-culture-infective dose (CCID₅₀). MT-4 cells were suspended in culture medium at 1×10^5 cells mL⁻¹ and infected with HIV at a multiplicity of infection of 0.02. Immediately after viral infection, 100 μ L of the cell suspension was placed in each well of a flat-bottomed microtiter tray containing various concentrations of the test compounds. Stock solutions of these compounds were dissolved in DMSO at 50 mm or higher. After incubation of virus-infected cells with the compounds at 37 °C for 4 days, the number of viable cells was determined using the MTT method. Compounds were tested in parallel for cytotoxic effects in uninfected MT-4 cells.

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- [1] D. W. Ludovici, B. L. De Corte, M. J. Kukla, H. Ye, C. Y. Ho, M. A. Lichenstein, R. W. Kavash, K. Andries, M. P. de Béthune, H. Azijn, R. Pauwels, P. J. Lewi, J. Heeres, L. M. H. Koymans, M. R. de Jonge, K. J. A. Van Aken, F. F. D. Daeyaert, K. Das, E. Arnold, P. A. J. Janssen, *Bioorg. Med. Chem. Lett.* 2001, 11, 2235–2239.
- [2] S. G. Sarafianos, K. Das, S. H. Hughes, E. Arnold, Curr. Opin. Struct. Biol. 2004, 14, 716–730.
- [3] K. Das, P. J. Lewi, S. H. Hughes, E. Arnold, Prog. Biophys. Mol. Biol. 2005, 88, 209–231.
- [4] B. L. De Corte, J. Med. Chem. 2005, 48, 1689-1696.
- [5] J. Heeres, P. J. Lewi, Adv. Antiviral Drug Des. 2007, 5, 213-242.
- [6] E. De Clercq, Nat. Rev. Drug Discovery 2007, 6, 1001-1018.
- [7] J. Heeres, M. R. de Jonge, L. M. H. Koymans, F. F. D. Daeyaert, M. Vinkers, K. J. A. Van Aken, E. Arnold, K. Das, A. Kilonda, G. J. Hoornaert, F. Compernolle, M. Cegla, R. A. Azzam, K. Andries, M. P. de Béthune, H. Azijn, R. Pauwels, P. J. Lewi, P. A. J. Janssen, J. Med. Chem. 2005, 48, 1910–1918.
- [8] J. Guillemont, E. Pasquier, P. Palandjian, D. Vernier, S. Gaurrand, P. J. Lewi, J. Heeres, M. R. de Jonge, L. M. H. Koymans, F. F. D. Daeyaert, M. H. Vinkers, E. Arnold, K. Das, R. Pauwels, K. Andries, M.-P. de Béthune, E. Bettens, K. Hertogs, P. Wigerinck, P. Timmerman, P. A. J. Janssen, J. Med. Chem. 2005, 48, 2072–2079.
- [9] C. Mordant, B. Schmitt, E. Pasquier, C. Demestre, L. Queguiner, C. Masungi, A. Peeters, L. Smeulders, E. Bettern, K. Hertogs, J. Heeres, P. Lewi, J. Guillemont, Eur. J. Med. Chem. 2007, 42, 567–579.
- [10] R. Haubrich, S. Gubernick, U. Yasothan, P. Kirkpatrick, Nat. Rev. Drug Discovery 2008, 7, 287–288.
- [11] G. Meng, F. E. Chen, E. De Clercq, J. Balzarini, C. Pannecouque, Chem. Pharm. Bull. 2003, 51, 779–789.
- [12] a) L. Ji, F. E. Chen, E. De Clercq, J. Balzarini, C. Pannecouque, J. Med. Chem. 2007, 50, 1778–1786; b) R. Cancio, A. Mai, D. Rotili, M. Artico, G. Sbardella, I. Clotet-Codina, J. A. Esté, E. Crespan, S. Zanoli, U. Hübscher, S. Spadari, G. Maga, ChemMedChem 2007, 2, 445–448; c) M. Radi, C. Falciani, L. Contemori, E. Petricci, G. Maga, A. Samuele, S. Zanoli, M. Terrazas, M. Castria, A. Togninelli, J. A. Esté, I. Clotet-Codina, M. Armand-Ugón, M. Botta, ChemMedChem 2008, 3, 573–593; d) A. Samuele, M. Facchini, D. Rotili, A. Mai, M. Artico, M. Armand-Ugón, J. A. Esté, G. Maga, ChemMedChem 2008, 3, 1412–1418.
- [13] a) Y. Z. Xiong, F. E. Chen, J. Balzarini, E. De Clercq, E. C. Pannecouque, Eur. J. Med. Chem. 2008, 43, 1230–1236; b) Y. Z. Xiong, F. E. Chen, X. Q. Feng, Acta Chim. Sin. 2006, 64, 1627–1630.
- [14] G. Sbardella, S. Bartolini, S. Castellano, M. Artico, N. Paesano, D. Rotili, C. Spadafora, A. Mai, ChemMedChem 2006, 1, 1073–1080.
- [15] a) Y. H. Liang, F. E. Chen, Eur. J. Med. Chem. 2008, DOI: 10.1016/j.ejmech.2008.03.021; b) Y. H. Liang, F. E. Chen, Drug Discovery Ther. 2007, 1, 57–60.
- [16] D. Roy, S. Ghosh, B. C. Guha, Arch. Biochem. Biophys. 1961, 92, 366–372.
- [17] R. Pauwels, J. Balzarini, M. Baba, R. Snoeck, D. Schols, P. Herdewijn, J. Desmyter, E. De Clercq, J. Virol. Methods 1988, 20, 309–321.
- [18] C. Pannecouque, D. Daelemans, E. De Clercq, Nat. Protoc. 2008, 3, 427–434.
- [19] D. S. Goodsell, G. M. Morris, A. J. Olson, J. Mol. Recognit. 1996, 9, 1-5.
- [20] K. Das, J. D. Bauman, A. D. Clark, Jr., Y. V. Frenkel, P. J. Lewi, A. J. Shatkin, S. H. Hughes, E. Arnold, *Proc. Natl. Acad. Sci. USA* 2008, 105, 1466–1471.
- [21] K. Das, A. D. Clark, Jr., P. J. Lewi, J. Heeres, M. R. de Jonge, L. M. H. Koymans, M. Vinkers, F. Daeyaert, D. W. Ludovici, M. J. Kukla, B. De Corte, R. W. Kavash, C. Y. Ho, H. Ye, M. A. Lichtenstein, K. Andries, R. Pauwels, M. P. de Béthune, P. L. Boyer, P. Clark, S. H. Hughes, P. A. J. Janssen, E. Arnold, J. Med. Chem. 2004, 47, 2550–2560.

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