Efficient Synthesis and Docking Analysis of Selective CDK9 Inhibitor NVP-2

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Abstract

NVP-2 (1), a potent and selective inhibitor of cyclin-dependent kinase 9 (CDK9), showed potent antitumor activity in preclinical studies. In this work, we designed and adopted a convergent synthetic route to efficiently synthesize NVP-2 (1). The key intermediate (7) was synthesized from malononitrile (2) and 1-bromo-2-(2-bromoethoxy)ethane (3) by successive cyclization, reduction, nucleophilic substitution with 2-bromo-6-fluoropyridine, and Suzuki–Miyaura reaction with (5-chloro-2-fluoropyridin-4-yl)boronic acid. Another key intermediate (11) was synthesized from (S)-1-methoxypropan-2-ol (8) by reaction with TsCl, electrophilic substitution reaction with tert-butyl ((1r,4r)-4-amino-cyclohexyl)carbamate, and then by deprotection of Boc. Finally, a substitution reaction by the key intermediates (7) and (11) to afford the target product NVP-2 (1). The reaction conditions of the whole synthesis process were simple and mild, free of harsh conditions such as the microwave reaction and dangerous reagents in the original patent, and realized the efficient synthesis of NVP-2. In addition, we analyzed the binding mode of NVP-2 in the active pocket of CDK9 to provide reasonable design ideas for subsequent discovery of novel CDK9 inhibitors.

Keywords

► NVP-2
► CDK9 inhibitor
► efficient synthesis
► docking analysis

# These authors contributed equally to this work.

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Introduction

Cyclin-dependent kinase 9 (CDK9) mainly regulates gene transcription elongation and messenger RNA (mRNA) maturation, and has emerged as an attractive therapeutic target for many malignant tumors, including hematological malignancies and solid tumors.\(^1\)–\(^5\) Many CDK9 inhibitors have been developed or are currently being in various stages of clinical development for cancer treatment.\(^6\)–\(^10\) NVP-2 (1), chemical name “4-(((5′-chloro-2′-(((1R,4R)-4-(((R)-1-methoxypropan-2-yl)amino)cyclohexyl)amino)-2,4′-bipyridin-6-yl)amino)methyl)tetrahydro-2H-pyran-4-carbonitrile,” is a novel selective ATP-competitive CDK9 inhibitor developed by Novartis in 2011. NVP-2 displays a very potent CDK9/CycT inhibitory activity with an IC50 value of 0.51 nmol/L, and more than 1,000-fold selectivity over CDK1/CDK2/CycA, and CDK16/CycY.\(^11\) In 2017, Winter et al found that NVP-2 significantly inhibited the proliferation of leukemia cells such as Jurkat, SKW-3, and SUP-T11, and induced the apoptosis of MOLT4 leukemia cells, suggesting the antitumor effect of NVP-2 in vitro.\(^11\) NVP-2 may become a promising antitumor agent to enter clinical research.

To date, a few synthesis routes of NVP-2 and its key intermediates have been reported. However, there are some shortcomings, such as harsh reaction conditions, low yields, side reactions, and potential safety hazards, in the reported synthetic route (Fig. 1).

The synthetic route of key intermediate 7 has been optimized and improved on the basis of comprehensive relevant literature methods:\(^12,13\) tetrahydro-4H-pyran-4,4-dicarbonitrile (4) was synthesized from inexpensive and commercially available malononitrile (2) and 2,2′-dibromodiethyl ether (3) as starting materials under milder conditions. Intermediate 4 was readily reduced to 4-(aminomethyl)tetrahydro-2H-pyran-4-carbonitrile (5) using the mild reducing agent NaBH₄. Nucleophilic substitution of 2-bromo-6-fluoropyridine by intermediate 5 occurs at high temperatures to give 4-(((6-bromopyridin-2-yl)amino)methyl)tetrahydro-2H-pyran-4-carbonitrile (6) with a high yield. Intermediate 6 and (5-chloro-2-fluoropyridin-4-yl)boric acid under Suzuki–Miyaura reaction conditions gave the key intermediate 4-(((5′-chloro-2′-fluoro-[2,4′-bipyridin]-6-yl)amino)methyl) tetrahydro-2H-pyran-4-carbonitrile (7). The synthetic route is shown in Scheme 1. In the improved synthetic route, we decreased the reaction temperature and appropriately extended the reaction time to efficiently obtain intermediate 4, thus overcoming the side reactions caused by the excessively high temperature. In the following step to synthesize intermediate 5, we could evade the formation of byproducts and improve the yield by slightly heating the reaction. Furthermore, in the Suzuki–Miyaura reaction, the harsh reaction conditions (high temperatures and microwave reactor) were used in the patent, which are not suitable for industrial manufacturing. We were able to avoid these harsh reaction conditions by changing the reaction solvent and employing an oil bath to effectively synthesize the key intermediate 7.

On the basis of the literature route, the synthetic route of key intermediate 11 is optimized and improved, shown in Scheme 2: inexpensive, readily available (S)-1-methoxy propan-2-ol (8) was used as the starting material, and reacted with p-toluenesulfonyl chloride to efficiently obtain (S)-1-methoxy-2-propyl-p-toluensulfonate (9). The mixture of intermediate (9) and tert-butyl ((1R,4R)-4-amino-cyclohexyl)carbamate was heated at reflux to efficiently

![Fig. 1](image-url) The reported synthetic route to NVP-2.\(^12,13\)
obtain the intermediate tert-butyl ((1R,4R)-4-((1R)-1-methoxypropan-2-yl)amino)cyclohexyl)carbamate (10). Intermediate 10 was quickly reacted with hydrochloric acid solution to deprotect the Boc group, and gave (1R,4R)-N1-((1R)-1-methoxypropan-2-yl)cyclohexane-1,4-diamine (11) hydrochloride salt. Intermediate 11 and the above key intermediate 7 undergo substitution reaction at a higher temperature to obtain the final product NVP-2. In the synthetic route in the literatures, trans-cyclohexanediamine was directly reacted with intermediate 9 at high temperature to obtain intermediate 11. However, there are some disadvantages, such as byproduct formation, difficulties at purification, and low yield in this step. Also, the hazardous chemical reagent NaH was utilized in the synthesis of intermediate 9 in the literatures, posing significant safety concerns in industrial manufacturing. In our improved synthetic route, the raw material tert-butyl ((1R,4R)-4-aminocyclohexyl)carbamate was directly reacted with intermediate 9 at a mild temperature to efficiently obtain intermediate 10 (65% yield), which quickly deprotected the Boc group in the acid solution to obtain key intermediate 11 (95% yield). By our improved synthetic route, we can efficiently obtain the pure key intermediate 11. In addition, we used pyridine as the base instead of the dangerous reagent NaH, and greatly shortened the reaction time to efficiently synthesize the intermediate 9 with a high yield of 90%. In the last step to synthesize the NVP-2, the addition of inorganic base potassium carbonate could reduce the reaction time and increase the yield. During the entire NVP-2 reaction process, the whole operation is simple, efficient and safe, and apt to industrial production.

Docking Analysis
NVP-2 competitively binds to the ATP active pocket of CDK9 protein to exert potent CDK9 inhibitory activity. However, the binding mode between NVP-2 and CDK9 protein has not been reported. Revealing and predicting the binding mode between NVP-2 and CDK9 protein could provide reasonable design ideas for the discovery of novel CDK9 inhibitors. Therefore, we applied a computer docking technology to simulate the interaction between NVP-2 and CDK9 protein. The predicted binding mode of NVP-2 in the ATP active pocket of CDK9 protein is shown in Fig. 2. In the ATP active pocket of CDK9 protein (Fig. 2A, B), it contains a very important hinge region and some hydrophobic pockets. In the hinge region, Cys106 residue can form vital hydrogen bonds with many small-molecule CDK9 inhibitors, including flavopiridol, variloxins, and others. The weakening of...
inhibitors based on the structure of NVP-2. The above docking analysis provided valuable clues for modification and discovery of novel CDK9 inhibitors based on the structure of NVP-2.

**Experimental Section**

**Chemistry**

Unless otherwise noted, all solvents and reagents were commercially available (Bidepharm, J&K Scientific, Sino-pharm) and used without further purification. All reactions were monitored by thin-layer chromatography on 0.25 mm silica gel plates (60 GF-254) and visualized with UV light, or iodine vapor. $^1$H NMR and $^{13}$C NMR were generated in DMSO-$d_6$ or CDCl$_3$ on Varian Mercury 400, 500, or 600 MHz NMR spectrometers. Chemical shifts were reported in parts per million (ppm). Multiplicity of $^1$H NMR signals was reported as single (s), double (d), triplet (t), quartet (q), and multiplet (m). ESI-MS (electrospray ionization mass spectra) were determined on an API 4000 spectrometer.

**Synthesis of Dihydro-2H-pyran-4,4(3H)-dicarbonitrile (4)**

A mixture of malononitrile (2, 7.0 g, 106 mmol), 1-bromo-2-(2-bromoethoxy)ethane (3, 24.57 g, 106 mmol) in DMF (100 mL) was stirred in an ice bath for 20 minutes. Then to the reaction was added DBU (1,8-diazabicyclo[5.4.0]undec-7-ene) (32.26 g, 212 mmol) dropwise. The reaction temperature was raised to 70°C and stirred for 3 hours. Then, the reaction was cooled to room temperature (r.t.), diluted with ethyl acetate, extracted with water and brine (150 mL $\times$ 3) until DMF was removed, and dried over anhydrous sodium sulfate. The organic phase was filtered and concentrated under reduced pressure to give a light brown solid (4, 12 g); yield: 83.18%. $^1$H NMR (600 MHz, DMSO-$d_6$) $\delta$ 3.73–3.67 (m, 4H), 2.32–2.26 (m, 4H).

**Synthesis of 4-(Aminomethyl)tetrahydro-2H-pyran-4-carbonitrile (5)**

To the solution of intermediate (4, 1.0 g, 7.34 mmol) in ethanol (35 mL) was slowly added sodium borohydride (0.83 g, 22.0 mmol) under ice bath conditions. The reaction mixture was removed from the ice bath and stirred at 60°C for 4 hours. The mixture was concentrated under reduced pressure and the resulting residue was diluted with ethyl acetate (100 mL), then washed with water and brine (150 mL $\times$ 3). The organic phase was dried over anhydrous sodium sulfate, filtered, and concentrated to dryness under vacuum to give a crude product (5), which was used directly in the next step. Low-resolution mass spectrometry (LRMS) m/z calcd. for C$_7$H$_{12}$N$_2$O $\left[M + H^+\right]^+$ 141.09, found: 141.

**Synthesis of 4-(((6-Bromo-pyridin-2-yl)amino)methyl)tetrahydro-2H-pyran-4-carboxylic carbonitrile (6)**

To a solution of intermediate (5, 1.35 g, 9.61 mmol) in DMSO (10 mL) was consistently added 2-bromo-6-fluoropyridine (1.41 g, 8.01 mmol) and triethylamine (2.43 g, 20.9 mmol). The temperature of the oil bath was raised to 110°C, and the reaction was refluxed for 18 hours. Upon completion of the reaction, the reaction mixture cooled to ambient temperature,
diluted with ethyl acetate, and washed with water and brine (150 mL × 3). After DMSO was washed and removed from the system, the organic layer was dried with anhydrous sodium sulfate, filtered, and concentrated under reduced pressure. The crude product was purified by flash column chromatography to give a light-yellow solid (6, 1.58 g); yield: 66.6%. LRMS m/z calc. for C_{17}H_{18}BrN_{3}O \[ M^+ \] 346.9. 1H NMR (400 MHz, CDCl₃) δ 7.24 (t, J = 7.8 Hz, 1H), 6.78 (d, J = 7.5 Hz, 1H), 6.39 (d, J = 8.2 Hz, 1H), 4.84 (t, J = 6.8 Hz, 1H), 4.00 (ddd, J = 12.4, 4.4, 2.2 Hz, 2H), 3.72–3.65 (m, 4H), 1.88 (dq, J = 13.8, 2.4 Hz, 2H), 1.77 (ddd, J = 13.8, 11.7, 4.4 Hz, 2H).

Synthesis of 4-(((5′-Chloro-2′-fluoro-[2,4′-bipyridyl]-6-yl) amino)methyl)tetrahydro-2H-pyrano-4-carbonitrile (7)

A mixture of 6 (1.0 g, 3.38 mmol), 5-chloro-2-fluoropyridin-4-yl-boronic acid (0.888 g, 5.06 mmol), Pd(dppf)Cl₂ (0.247 g, 0.38 mmol), and THF/H₂O (2:1 [v/v], 51 mL) was stirred in an ice bath for 20 minutes, and to the reaction mixture was sequentially added (20 mL) was stirred in an ice bath for 20 minutes, and to the reaction mixture was cooled to ambient temperature and concentrated in vacuo. The crude product was dissolved in ethyl acetate, was cooled to r.t., diluted with water, and extracted with ethyl acetate. The organic layer was washed with brine (100 mL × 3), dried over anhydrous sodium sulfate, filtered, and concentrated under reduced pressure. The resulting crude material was purified by flash chromatography to give a pale-yellow solid (10, 0.76 g); yield: 65.0%. LRMS m/z calc. for C₁₅H₁₇N₃O₂ \[ M^+ \] 287.23, found: 287.1. 1H NMR (400 MHz, CDCl₃) δ 4.38 (s, 1H), 3.41 (s, 1H), 3.34 (s, 3H), 3.29 (dd, J = 9.2, 4.5 Hz, 1H), 3.23 (dd, J = 9.3, 7.0 Hz, 1H), 3.01 (td, J = 6.7, 4.7 Hz, 1H), 2.51 (t, J = 11.2, 3.9 Hz, 1H), 2.07–1.97 (m, 3H), 1.92 (dp, J = 9.3, 3.3 Hz, 1H), 1.44 (s, 9H), 1.24–1.09 (m, 4H), 1.03 (d, J = 6.4 Hz, 3H).

Synthesis of (1r,4R)-N^1-((R)-1-Methoxypropan-2-yl)cyclohexane-1,4-diamine (11)

To the solvent of intermediate 10 (1.0 g, 3.49 mmol) in DCM (5 mL) was slowly added 4 mol/L hydrochloric acid in dioxane (10 mL). Then the reaction was stirred at r.t. for 4 hours. The reaction mixture was concentrated in vacuo to give a white solid (11, 0.62 g); yield: 95.0%. LRMS m/z calc. for C₁₀H₂₂N₂O \[ M^+ \] 187.30, found: 187.1. 1H NMR (400 MHz, CDCl₃) δ 3.34 (s, 3H), 3.28 (dd, J = 9.2, 4.5 Hz, 1H), 3.22 (dd, J = 9.2, 6.9 Hz, 1H), 3.00 (td, J = 6.6, 4.7 Hz, 1H), 2.66 (ddt, J = 14.6, 9.5, 3.8 Hz, 1H), 2.49 (t, J = 10.9, 3.9 Hz, 1H), 1.97 (dt, J = 12.3, 3.2 Hz, 1H), 1.93–1.83 (m, 3H), 1.26 (s, 2H), 1.17–1.12 (m, 2H), 1.02 (d, J = 6.4 Hz, 3H).

Synthesis of NVP-2 (1)

A mixture of intermediate 7 (0.5 g, 1.44 mmol), intermediate 11 (0.223 g, 1.73 mmol), and K₂CO₃ (0.598 g, 4.33 mmol) in DMSO (20 mL) was stirred at 110°C for 16 hours. When TLC monitoring showed the starting material was nearly completely consumed in the reaction, then the reaction was cooled to r.t., diluted with ethyl acetate, and washed with water (50 mL × 3). The organic phase was dried with anhydrous sodium sulfate, filtered, and concentrated under reduced pressure. The resulting crude material was purified by flash chromatography to obtain NVP-2 (1, 0.21 g) as a gray solid; yield: 28.4%. LRMS m/z calc. for C_{26}H_{26}ClN_{13}O_{3} \[ M^+ \] 513.27, found: 512.8. 1H NMR (500 MHz, DMSO-d₆) δ 8.02 (s, 1H), 7.50 (dd, J = 8.4, 7.2 Hz, 1H), 7.08 (t, J = 6.5 Hz, 1H), 6.78 (d, J = 7.2 Hz, 1H), 6.67 (d, J = 8.4 Hz, 1H), 6.61 (d, J = 11.7 Hz, 2H), 3.90 (dd, J = 12.1, 4.5, 2.1 Hz, 2H), 3.67 (d, J = 6.4 Hz, 2H), 3.62–3.53 (m, 1H), 3.48 (td, J = 12.1, 2.0 Hz, 2H), 3.25 (s, 3H), 3.21–3.18 (m, 1H), 3.14 (dd, J = 9.2, 5.5 Hz, 1H), 2.92 (p, J = 6.1 Hz, 1H), 2.50–2.45 (m, 1H), 1.97–1.83 (m, 5H), 1.68 (dd, J = 13.7, 11.8, 4.5 Hz, 2H), 1.26–1.02 (m, 5H), 0.94 (d, J = 6.3 Hz, 3H). 13C NMR (126 MHz, DMSO-d₆) δ 158.26, 157.24, 152.31, 147.07, 146.75, 137.06, 122.45, 114.97, 112.40, 109.25, 108.58, 77.12, 63.85, 58.12, 52.76, 49.38, 48.47, 47.19, 38.62, 32.71, 32.41, 31.88, 31.32, 31.16, 18.15.

Docking Simulation

The binding mode between NVP-2 and the CDK9 active pocket was completed by the conventional computer docking software Schrödinger-Maestro (https://www.schrodinger.com/products/matego, version 11.0, Schrödinger software package, New York,
The structural optimization of CDK9 protein (PDB number: 4BCF) was completed by the Protein Preparation Wizard sector in Maestro. The structure characterization of CDK9 protein was performed by adding hydrogens, creating disulfide bonds, filling in missing side chains, deleting water, restrained minimization, and other optimization operations (other parameters are kept unchanged). Subsequently, the receptor grid box (10 × 10 × 10 Å) was obtained based on the optimized structure of CDK9 protein. Continue to use LigPrep in Maestro to process the NVP-2 structure to generate various three-dimensional molecular configurations, and then used the Ligand Docking sector to complete the docking simulation between NVP-2 and CDK9 protein.

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Conflict of Interest
All authors declare no conflict of interest.

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