Efficient and Facile Synthesis of Chromenopyrano[2,3-b]pyridine Derivatives Catalyzed by Sodium Carbonate

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Abstract

In this research, a number of new and known chromenopyrano[2,3-b]pyridine derivatives have been prepared. Initially, according to the reported procedure, pyrano[2,3-c]chromene derivatives were synthesized by the reaction between 4-hydroxycoumarin, aromatic aldehydes and malononitrile using silica sodium carbonate (SSC) as the catalyst. Next, the prepared pyrano[2,3-c]chromenes were reacted by dimethyl acetylenedicarboxylate (DMAD) or cyclohexanone in the presence of sodium carbonate to produce chromenopyrano[2,3-b]pyridine derivatives. The presented protocol avoids the use of expensive catalysts and gives useful potentially bioactive heterocycles in excellent to high yields.

Keywords: Chromenopyrano[2,3-b]pyridine; pyrano[2,3-c]chromene, dimethyl acetylenedicarboxylate, cyclohexanone

1. Introduction

Heterocycles as the most prevalent organic compounds are present in various drugs, natural products, vitamins and bioactive compounds.1-3 They have been found to be useful as anti-HIV,4 anti-tumor,5-6 anti-inflammatory,7-9 anti-malarial,10,11 anti-depressant, anti-microbial, anti-bacterial and anti-fungal agents.12 Most heterocycles are used in materials science such as fluorescent sensors, dyes, data storage, plastics, illuminators and analytical reagents.5,13 Also, they are applied as important intermediates for the synthesis of medicinal compounds.14 Nitrogen-containing six-membered heterocycles, due to their high biological activity are of interest to both medicinal chemists and biochemists.15,16 Among them, pyridine and its derivatives play the most fundamental structural role in many natural compounds and medicinally beneficial molecules.17,18 Owing to the great variety of biologically active pyridines, it is not surprising that the pyridine ring system has become a vital basic component in many pharmaceutical agents. Some of the pyridine-derived drugs are trademarks of Reyataz and Gleevec drugs that are prescribed for HIV and chronic anemia, respectively. Some natural alkaloid products based on pyridine are nicotine and niacin derivatives.19 In particular, condensed pyridines are known for their several biological activities. For example, pyranopyridines have been proven to be the most active anti-tumor heterocyclic systems with activity against various filamentous tumors and specific activity against lung and ovarian cancer cells and antimicrobial action.20,21 Moreover, pyrazolo-[3,4-b]pyridine derivatives are an important class of fused pyridines with a broad spectrum of biological activities that find widespread use in the pharmaceutical industries.22 Another class of fused pyridine derivatives is thiazolo[4,5-b]pyridin, which have anti-inflammatory and antimicrobial activity against human and veterinary pathogens. Also, some of these compounds have antifungal and anti-tumor activity.23,24

Coumarin-thiazole scaffolds are used as fluorescence probes for staining and imaging of DNA and to study the biological function of cell membranes.25,26 Other compounds fused to the coumarin ring, such as pyranochromenes as the most imperative fused polycyclic heterocycles have gained attention for their extensive occurrence in important pharmaceutical drugs.27 Many of pyranochromene derivatives exhibit significant biological activity, such as excellent antimicrobial potency, spasmolytic, anticoagulant, diuretic, cytotoxic, antituberculosis, anticancer, and antianaphylactic activities.28-30 Some of these compounds have found the use for the treatment of neurodegenerative diseases, including Parkinson’s disease, Down’s syndrome, Alzheimer’s disease and AIDS associated dementia.31 Furthermore, pyranochromene derivatives have been applied as the key intermediates for the preparation of thioxo-imidazolidinedione, dithioxodiazetidine and Schiff’s bases.32
Considering the above reports and in connection with our program on the synthesis of polycyclic compounds, we present in this paper an efficient and environmentally benign strategy for the synthesis of pyranochromene fused with pyridine derivatives. For this purpose, initially, pyrano[2,3-c]chromenes were prepared via the reaction between 4-hydroxycumarin (1), aromatic aldehydes 2 and malononitrile (3) using silica sodium carbonate (SSC) (Scheme 1). Subsequently, the synthesized pyrano[2,3-c]chromenes were reacted by dimethyl acetylenedicarboxylate (DMAD, 5) or cyclohexanone (6) to yield chromenopyrano[2,3-b]pyridine derivatives 7 and 8 (Scheme 2).

Scheme 1. SSC-catalyzed synthesis of pyranochromenes 4.

Scheme 2. Synthesis of chromenopyrano[2,3-b]pyridine derivatives 7 and 8 in the presence of Na2CO3 as the catalyst.

2. Results and Discussion

First, pyrano[2,3-c]chromenes 4 were prepared and identified according to the mentioned method (Table 1). In the following, in order to optimize the conditions, the reaction between pyranochromene 4a and DMAD was selected as a model system. The reaction was not completed in the absence of a catalyst at room temperature. Next, the model reaction was performed in the presence of 5 mol% of Na2CO3 in various solvents such as acetone, CH2Cl2, EtOAc, DMF and DMSO. As can be seen in Table 2, the best result was obtained by performing the reaction mixture in DMSO (100 °C) to yield product 7a. Next, we evaluated the required amount of the catalyst for this transformation. When 10 mol% of Na2CO3 was used, the reaction efficiently proceeded and was complete in shorter reaction time. By further increasing the catalyst amount no appreciable improvement in the product yield and reaction time was observed. Various bases were screened for their efficiency in this reaction. We obtained the best yield of 7a when the reaction was performed by Na2CO3. Also, low temperatures led to the reaction product in a very low yield. In short, according to the obtained results, the best yield was achieved in DMSO at 100 °C in the presence of 10 mol% of Na2CO3 (Table 2, Entry 8). In view of the success of the above reaction and having established the optimal conditions, we then investigated the scope and general applicability of this methodology by using different pyranochromenes and the results are given in Table 3. The structures of the synthesized compounds 7 were deduced from their elemental analysis, IR, 1H and 13C NMR spectroscopy. Subsequently, by employing this method, a number of chromenopyrano[2,3-b]pyridines 8 were produced via the reaction between pyrano[2,3-c]chromenes 4 and cyclohexanone (6) under the optimized reaction conditions. In the majority of cases, the reactions performed cleanly and the desirable products 8 were formed in high yield.
Table 1. Synthesis of pyrano[2,3-c]chromenes 4 using SSC catalyst.

| Entry | Ar            | Product | Yield\(^a\) (%) | Mp [Ref.] (°C) |
|-------|---------------|---------|-----------------|----------------|
| 4a    | C\(_6\)H\(_5\) | 83      | 260–262 [261–263]\(^{28}\) |
| 4b    | 4-Cl-C\(_6\)H\(_4\) | 85      | 262–264 [262–264]\(^{28}\) |
| 4c    | 4-Br-C\(_6\)H\(_4\) | 80      | 245–247 [247–249]\(^{41}\) |
| 4d    | 3-NO\(_2\)-C\(_6\)H\(_4\) | 85      | 264–265 [263–265]\(^{28}\) |
| 4e    | 4-NO\(_2\)-C\(_6\)H\(_4\) | 90      | 259–261 [260–262]\(^{28}\) |
| 4f    | 3-CH\(_3\)-C\(_6\)H\(_4\) | 90      | 255–257\(^b\) |
| 4g    | 4-F-C\(_6\)H\(_4\) | 93      | 258–260 [260–262]\(^{41}\) |
| 4h    | 4-CH\(_3\)-C\(_6\)H\(_4\) | 87      | 250–252 [255–257]\(^{28}\) |
| 4i    | 2-Cl-C\(_6\)H\(_4\) | 85      | 271–272 [269–271]\(^{28}\) |
| 4j    | 4-OCH\(_3\)-C\(_6\)H\(_4\) | 80      | 243–245 [246–248]\(^{28}\) |
| 4k    | 2,4-Cl\(_2\)-C\(_6\)H\(_4\) | 83      | 259–260 [259–260]\(^{28}\) |

\(^a\) Isolated yield.
Table 2. Synthetic results of 7a under different reaction conditions.

| Entry | Catalyst (mol %) | Solvent/Temp. (°C) | Time (h) | Yield* (%) |
|-------|------------------|---------------------|----------|------------|
| 1     | None             | None/25             | 24       | 5          |
| 2     | Na₂CO₃ (5)       | EtOAc/reflux        | 5        | 40         |
| 3     | Na₂CO₃ (5)       | DMSO/100            | 5        | 50         |
| 4     | Na₂CO₃ (5)       | CH₂Cl₂/reflux       | 5        | 30         |
| 5     | Na₂CO₃ (5)       | Acetone/reflux      | 5        | 40         |
| 6     | Na₂CO₃ (5)       | DMF/100             | 5        | 30         |
| 7     | Na₂CO₃ (1)       | DMSO/100            | 5        | 30         |
| 8     | Na₂CO₃ (10)      | DMSO/100            | 4        | 80         |
| 9     | Na₂CO₃ (15)      | DMSO/100            | 4        | 75         |
| 10    | Piperidine (10)  | DMSO/100            | 5        | 35         |
| 11    | NaOH (10)        | DMSO/100            | 5        | 42         |
| 12    | KOH (10)         | DMSO/100            | 5        | 45         |
| 13    | Na₂CO₃ (10)      | DMSO/50             | 4        | 37         |
| 14    | Na₂CO₃ (10)      | DMSO/70             | 4        | 50         |

* Isolated yield.

Table 3. Synthesis of chromenopyrano[2,3-b]pyridine derivatives 7 and 8 in the presence of Na₂CO₃ catalyst.

| Entry | Ar             | Product Time (h) | Yield* (%) | Mp [Ref.] (°C) |
|-------|----------------|-----------------|------------|----------------|
| 7a    | C₆H₅           | 4               | 80         | decomp. >300b  |
| 7b    | 4-Cl-C₆H₄      | 3               | 75         | decomp. >300b  |
| 7c    | 4-Br-C₆H₄      | 3               | 73         | 290–291b       |

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| Substituent     | Yields | Temperature | Decomposition Temperature |
|-----------------|--------|-------------|---------------------------|
| 7d 3-NO$_2$-C$_6$H$_4$ | 3.5    | 80          | decomp. >300$^b$          |
| 8a C$_6$H$_5$    | 7      | 85          | 267–269 (>260)$^{42}$     |
| 8b 4-NO$_2$-C$_6$H$_4$ | 6     | 82          | 270–272$^b$               |
| 8c 3-CH$_3$-C$_6$H$_4$ | 7     | 75          | 269–271$^b$               |
| 8d 4-F-C$_6$H$_4$  | 8      | 70          | decomp. >260 (>260)$^{42}$|
| 8e 4-CH$_3$-C$_6$H$_4$ | 6     | 90          | 265–268 (>260)$^{42}$     |
| 8f 3-NO$_2$-C$_6$H$_4$ | 5     | 90          | decomp. >260 (>260)$^{42}$|
| 8g 2-Cl-C$_6$H$_4$ | 5.5    | 80          | 259–261 (>260)$^{42}$     |
| 8h 4-OCH$_3$-C$_6$H$_4$ | 8     | 95          | 263–265 (>260)$^{42}$     |
yields and the nature of the Ar group appeared to have no notable effect on the reaction rate (Table 3).

We propose a mechanism for the Na2CO3-catalyzed synthesis of chromenopyranopyridine derivatives 7 (Scheme 3). Firstly, intermediate 9 is produced by the deprotonation of NH2 of pyranochromene in the presence of Na2CO3. Next, a Michael-type addition of NH – to dimethyl acetylenedicarboxylate creates adduct 10. The intramolecular cyclization of 10 gives adduct 11 which rearranges into the product 7. The synthesis of product 8 also can be visualized as proceeding through a condensation of intermediate 9 with cyclohexanone to produce 12. Subsequently, cyclization and then tautomerization of 12 gives the desired product 8 (Scheme 3).

3. Experimental

All chemicals and reagents were purchased from Fluka and Merck companies. The reaction progress was monitored by using TLC on silica gel polygram SIL G/UV254 plates. Reported melting points were determined by an electrothermal KSB1N apparatus. 1H NMR spectra were recorded in DMSO-d6 on a Bruker Avance Ultra Shield 400 MHz instrument spectrometer and 13C NMR spectra were recorded at 100 MHz. A Vario-El CHN instrument at the Isfahan Industrial University was used for the elemental analyses. IR spectra were obtained with a JASCO FT-IR/680 instrument spectrometer using KBr pellets.

Procedure for the Preparation of the SSC Catalyst

Silica sodium carbonate (SSC) was synthesized as a low-cost, recyclable and green catalyst in two steps according to the literature procedure.43 Initially, by adding thionyl chloride gradually to the silica gel, the silica chloride was prepared. Then, silica chloride and sodium bicarbonate were refluxed in hexane solvent for 24 hours. The solid product obtained, after washing and drying was used as a silica sodium carbonate catalyst.

General Procedure for the Synthesis of Pyranochromenes 4

SSC (0.1 mmol, 0.2 g) was added to a mixture of malononitrile, aryl aldehyde, and 4-hydroxycoumarin at 110 °C under solvent-free conditions. The reaction progress was monitored by TLC. After completion of the reaction, boiling EtOAc (10 mL) was added, and the catalyst was separated by filtration. To further purify the product, obtained powder was recrystallized from EtOH.28

Preparation of Chromenopyranopyridines 7

A solution of DMAD (1 mmol), compound 4 (1 mmol) and Na2CO3 (10 mol%) in DMSO (5 mL) was stirred at 100 °C for the appropriate time. The progress of the reaction was monitored by TLC (EtOAc/n-hexan). After completion of the reaction, the catalyst was separated by filtration and the solvent was evaporated under reduced pressure. The obtained products were purified by column chromatography.

Preparation of Chromenopyranopyridines 8

A mixture of compound 4 (1 mmol), cyclohexanone (5) (2 mmol) and Na2CO3 (10 mol%) in DMSO (5 mL) was stirred and heated at 100 °C for a required time. After termination of the reaction as demonstrated by TLC (EtOAc/n-hexan), the catalyst was separated by filtration. The solvent was removed and the crude powder 8 was purified by crystallization from EtOAc.

Dimethyl 8-Amino-6-oxo-7-phenyl-6H,7H-chromeno[3’,4’:5,6]pyranopyridine-9,10-dicarboxylate (7a). Yield: 80% (0.35 g), IR (KBr) (νmax cm−1) 3446, 2918, 1650, 1640, 1321. 1H NMR (400 MHz, DMSO-d6) δ 8.15 (s, 2H), 7.52–8.13 (m, 3H), 7.42-7.50 (m, 3H), 7.24–7.34 (m, 3H), 5.21 (s, 1H), 2.35 (s, 3H), 2.24 (s, 3H).

13CNMR (100 MHz, DMSO-d6) δ 160.16, 152.78, 151.24, 142.01, 132.81, 128.59, 124.63, 122.37, 117.23, 113.48, 100.22, 100.16, 77.35, 77.24, 77.04, 76.72, 53.22, 34.95, 30.98. Anal. Calcd for C25H18N2O7 C, 65.50; H, 3.96; N, 6.11. Found C, 65.53; H, 3.90; N, 6.18.
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Povzetek

V okviru te raziskave smo pripravili serijo novih in nekaj že znanih kromenopirano[2,3-b]piridinskih derivatov. Skladno z že objavljenimi postopki smo najprej z reakcijo med 4-hidroksikumarinom, aromatskimi aldehidi in malononitrilom ob prisotnosti silika natrijevega karbonata (SSC) kot katalizatorja pripravili pirano[2,3-c]kromenske derivate, ki smo jih v naslednji stopnji reagirali z dimetil acetilendikarboksilatom (DMAD) ali cikloheksanonom v prisotnosti natrijevega karbonata. Tako smo pripravili serijo kromenopirano[2,3-b]piridinskih derivatov. Za izvedbo našega protokola ne potrebujemo dragih katalizatorjev in vendar lahko potencialno bioaktivne heterociklične spojine pripravimo z odličnimi do visokimi izkoristki.