

A convenient synthesis of cyclopenta[b]pyridin-2,5-dione as a non-glycosidic cardiotonic agent

Nicolas Robert, Cécile Verrier, Christophe Hoarau, Sylvain Célanire,
and Francis Marsais*

UMR 6014 - CNRS, INSA and University of Rouen
IRCOF - INSA of Rouen BP 08 76131 Mont-Saint-Aignan Cédex, France
E-mail: francis.marsais@insa-rouen.fr

Dedicated to Professor Guy Quéguiner on the occasion of his 70th birthday

Abstract

A straightforward synthesis of cyclopenta[b]pyridin-2,5-dione is reported starting from the commercially available 2-bromo-6-methoxypyridine. The overall route consists in a first sequence of regioselective ortho lithiation and methoxycarbonylation followed by Heck vinylation, alkene reduction, cyclization and decarboxylation.

Keywords: Lithiation, Heck vinylation, pyridine, cyclopenta[b]pyridine-2,5-dione

Introduction

A recent pharmacological evaluation of various functionalized 2-pyridones as cardiotonic agents has revealed that the cyclopenta[b]pyridin-2,5-dione (**1**) displays a high activity rather similar to Milrinone (**2**) which is the most effective non glycosidic cardiotonic agent clinically used for the treatment of severe heart failure.¹ Cyclopenta[b]pyridin-2,5-dione (**1**) constitutes also an interesting tensor of pharmaceutics exemplified by the antibacterial product **5** and a building-block for the access to 2-cyclopenta[b]pyridin-5-one (**3**) as seco analogues of 8-azasteroids (**4**).²

Results and Discussion

Despite the fact that the cyclopenta[b]pyridin-2,5-dione (**1**) is gaining interest as biologically active compounds and valuable building-blocks only two methods of preparation could be found in the literature. The first synthesis of cyclopenta[b]pyridin-2,5-dione (**1**) was first reported in 1954 (6 steps synthesis and a 13 % overall yield).³ Mosti and his team published in 2003 a novel

synthetic route based upon a one-pot and two-step construction of the 2-pyridone ring from the cyclopenta-1,3-dione.¹ We recently described a novel synthesis of 6-methyl cyclopenta[*b*]pyridin-5-one (**8**) based on Heck vinylation of 2-bromo-6-methyl nicotinate (**6**) with methacrylate affording the pyridylacrylate intermediate **7**, alkene reduction and Dieckmann condensation as depicted in Scheme 1.⁴ We wish to report here our results on the application of the latter method to the preparation of the cyclopenta[*b*]pyridin-2,5-dione (**1**). Our retrosynthetic analysis suggests that 2-bromo-6-methoxynicotinate (**10**) could be a valuable precursor for this purpose (scheme 1). The pyridylacrylate **9** could be first prepared by Heck vinylation of bromopyridine **10**. The expected cyclopenta[*b*]pyridin-2,5-dione (**1**) would be then obtained by reduction of the alkene followed by a cyclization-decarboxylation sequence. The success of this novel approach mainly depends on the access to the unknown 2-bromo-6-methoxynicotinate **10**. Two possible routes could be designed: (i) the regioselective displacement of a bromine atom at position 2 of the methyl 2,6-dibromonicotinate (**11**) which could be readily prepared in two steps from the 2,6-dichloronicotinic acid by bromination and esterification⁷ or (ii), the regioselective methoxycarbonylation of the commercially available 2-bromo-6-methoxypyridine (**12**).

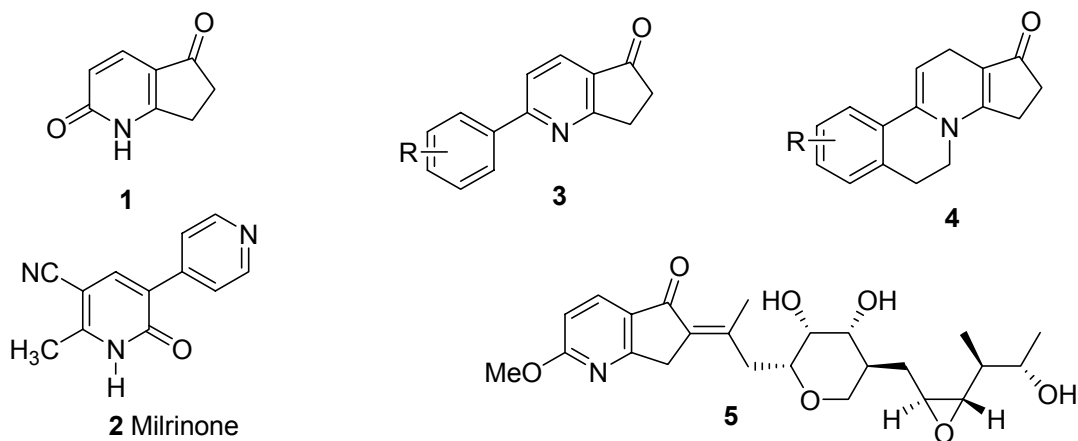
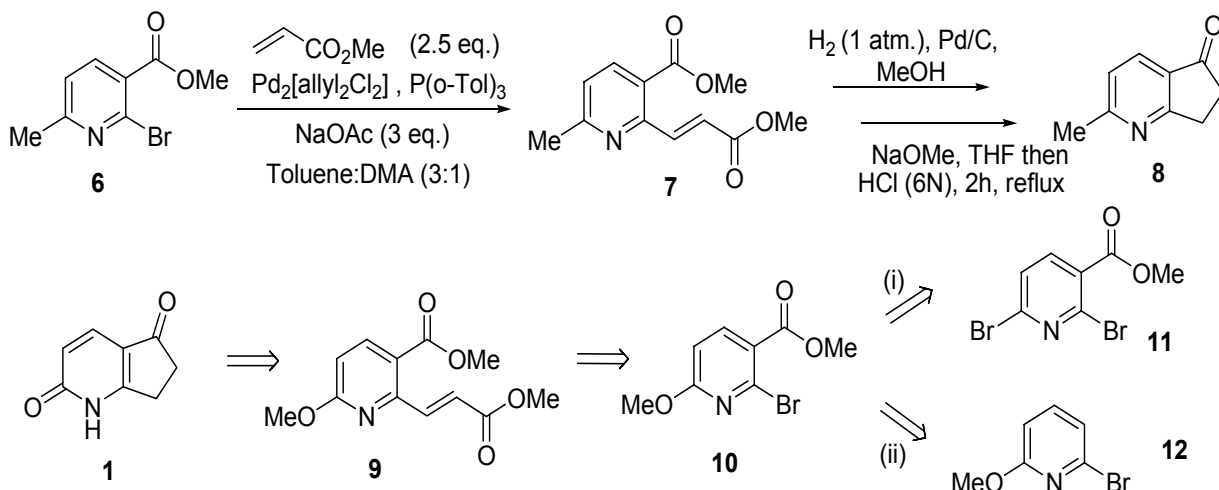
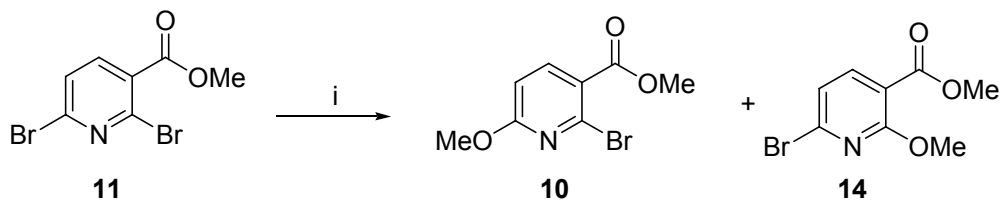


Figure 1



Scheme 1. Previously reported synthesis of 6-methylcyclopenta[b]pyridine-5-one (**8**) and retrosynthetic analysis of cyclopenta[b]pyridin-2,5-dione (**1**).

We first attempted to displace the bromine atom at position 2 of ethyl 2,6-dibromonicotinate (**11**) with sodium methoxide. Treatment of **11** with 1.5 equivalent of sodium methoxide was carried out in refluxing MeOH for 24h following the Hirokawa's protocol.⁵ A complete conversion of the starting material was observed and a mixture of 2- and 6- monosubstituted products **10** and **14** in a 7:3 ratio (¹H NMR) could be obtained in 75 % yield. Unfortunately, the two regioisomers **10** and **14** could not be separated by chromatography. Moreover, replacement of sodium methoxide by potassium methoxide also led to a 1:1 mixture regioisomers **10** and **14**.



Scheme 2. Reagents and conditions: (i) MeONa (1 equiv.), MeOH, reflux, 24h, 75%.

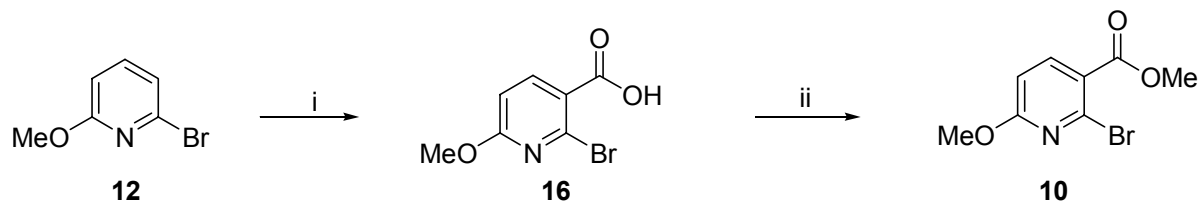
We then shifted to the second route based on the regioselective methoxycarbonylation at position 3 of 2-bromo-6-methoxy pyridine (**12**). To this purpose, we first examined the lithiation of 2-bromo-6-methoxy pyridine (**12**) by treatment with hard bases such as lithium amides in THF before quenching the lithio intermediates with D_2O (Table 1). A first set of lithiation experiments was achieved using 2,2',6,6'-tetramethylpiperidinyl-lithium (LTMP) at -78 °C (entries 1-3).

Table 1. Assays of regioselective lithiation of **12** with lithium amides

Entry	Base	Equiv.	T (°C)	15a (%)^a	15b (%)^a
1		1	-78°C	-	-
2	LTMP	2	-78	0	64
3		3	°C -78°C	10 15	80 -
4		2	-78	15	-
5	LDA	3	°C -78°C	15	-
6		2	-50	22	12
7		3	°C -50°C	20	11

^a The ratio was determined by ¹H NMR spectroscopy

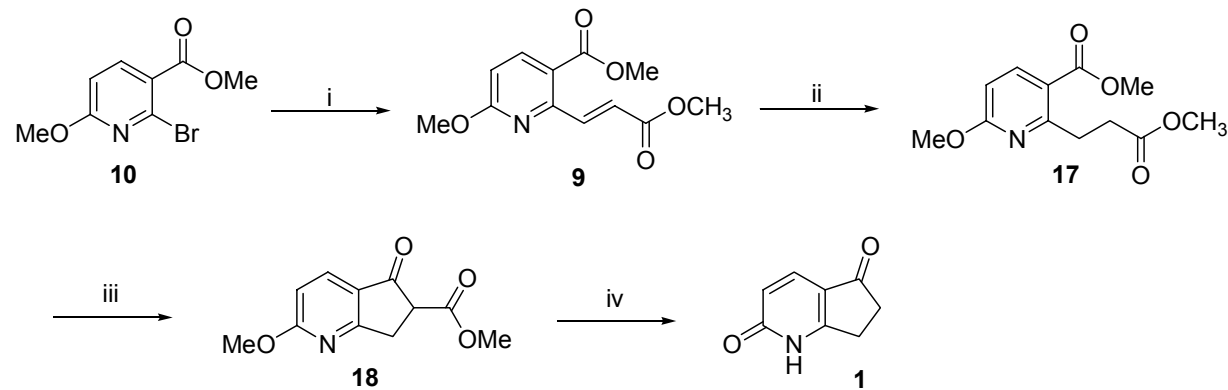
No deuterated product was obtained using 1 equivalent of LTMP whereas the 5-deuterated compound **15b** was selectively formed in 64% yield using 2 equivalents of LTMP. The starting material conversion could be significantly improved employing 3 equivalents of LTMP leading to a mixture of 3- and 5-deuterated products (**15a**, **15b**) in 2:8 ratio in favor of the **15b** isomer. The less hard lithium diisopropylamine (LDA) was also checked (entries 4-7). Treatment of **12** with 2 equivalents of LDA followed by D₂O trapping specifically provided the 3-deuterated product **15a** in 15 % yield (entry 4). This result could be related to the higher acidity of the proton at position 3. Surprisingly, we observed that the yield of 3-deuterated compound **15a** was not improved using 3 equivalents of LDA at the same temperature (entry 5). Moreover warming the 3-lithio anion from -78°C to -50 °C⁶ before trapping with D₂O afforded a mixture of **15a** and **15b** (entries 6,7).



Scheme 3. Reagents and conditions: (i) (a) LDA (2 equiv.), -78°C, THF, 1h, (b) solid carbon dioxide, (c) HCl (2M), 13 %; (ii) (a) (COCl)₂, DMFcat., (b) MeOH, 92%.

The 3-lithio anion formed by treatment of **12** with 2 equivalents of LDA at -78°C in THF (table 1-entry 4) did not react with methyl cyanoformate but could be trapped by carbon dioxide to give the 3-bromo-6-methoxynicotinic acid (**16**) after acidic treatment. Acid (**16**) was isolated in 13 % yield but the unreacted starting material **12** could be readily recovered and re-used. Finally **16** was obtained in 41 % overall yield after five lithiation–carboxylation sequences. Esterification of **16** gave the expected methyl 2-bromo-6-methoxy nicotinate (**10**) in 92 % yield.

Heck vinylation of **10** with methyl acrylate using the η^3 -allylpalladium chloride dimer with $P(o\text{-Tol})_3$ complex as catalyst in toluene and dimethylacetamide (DMA) as co-solvent⁴ provided the β -2-pyridyl acrylate **9** in an excellent 82% yield. The use of DMA is a crucial parameter as the vinylation of **10** failed without this co-solvent. Reduction of the alkene under soft conditions provided diester **17** which could be cyclized by a Dieckmann condensation with sodium methoxide to methyl cyclopenta[b]pyridine-5-one-6-carboxylate (**18**) in 70% overall yield. Finally, treatment with hydrochloric acid allows hydrolysis, decarboxylation and demethylation of **18** to give cyclopenta[c]pyridine-2,5-dione (**1**) in 84 % yield.



Scheme 4. Reagents and conditions: (i) methyl acrylate (2.5 equiv.), $Pd_2[allyl_2Cl_2]$, $P(o\text{-Tol})_3$, NaOAc (3 equiv.), toluene: DMA (3:1), 92%; (ii) H_2 (1 atm), Pd/C (10 mol %), MeOH, r.t., 2h, 93%; (iii) NaOMe, THF, reflux, 2h, 75%; (iv) HCl (5 M), reflux, 3h, 84%.

Conclusions

A convenient route to cyclopenta[b]pyridin-2,5-dione (**1**) is reported starting from methyl 2-bromo-6-methoxynicotinate (**10**) through a 4 steps synthesis, vinylation, alkene reduction and cyclization-decarboxylation, in 48% overall yield. Two routes were studied for the previous preparation of the parent methyl 2-bromo-6-methoxynicotinate (**10**). Regioselective displacement of the bromine atom of methyl 2,6-dibromonicotinate (**11**) by the sodium or potassium methoxide could not be applied leading to a mixture of regioisomers which could not be separated by chromatography. The second approach was based upon the regioselective

carboxylation of the commercially available 2-bromo-6-methoxypyridine (**12**) at position 3 of the pyridine nucleus. The regioselective lithiation-carboxylation and esterification of **12** at position 3 was achieved using LDA at -78°C in THF to give the expected methyl 2-bromo-6-methoxynicotinate (**10**) in 12 % yield in two steps.

Experimental Section

General Procedures. Tetrahydrofuran (THF), ether (Et_2O) were pre-dried with pellets of KOH and distilled over sodium benzophenone ketyl under Ar before use. CH_2Cl_2 , NEt_3 and toluene were distilled from CaH_2 . Methanol and ethanol were distilled from magnesium turnings; dimethylacetamide was distilled over 4 Å molecular sieves. For Flash chromatography, Merck silica gel (70-230 mesh) was used. The melting points were measured on a Kofler melting points apparatus and were not corrected. The ^1H NMR and ^{13}C NMR spectra were recorded with a Bruker Avance-300 spectrometer operating at 300 MHz. Commercially available starting materials were used without further purification. Infrared spectra were recorded on a Perkin-Elmer FT-IR 1650 spectrophotometer. Elemental analysis of compounds was carried out on a Carlo Erba 1160. Mass spectra were recorded on a JEOL JMS AX-500 spectrometer, in electronic impact (EI). The starting compound **12** is commercially available.

Preparation of 6-methoxy-2-bromonicotinate (**10**)

Methyl 2,6-dibromonicotinate (11**).** To a stirred solution of 2,6-dibromonicotinic acid⁷ (500 mg, 1.8 mmol) and 3 drops of DMF in dry CH_2Cl_2 (10 ml) was slowly added oxalyl chloride (172 μL , 2.0 mmol) at 0°C. The mixture was stirred at room temperature for 1h and solvents were removed *in vacuo*. To the crude product was added dry methanol (10 ml) at 0 °C and the resulting solution was stirred for 2 h at room temperature. Methanol was removed *in vacuo* and the crude solid was dissolved in CH_2Cl_2 (10 ml). The pH of the aqueous layer was then adjusted to 7 by adding aq. K_2CO_3 (2M). The separated organic layer was washed three times with water, dried (MgSO_4) and concentrated *in vacuo* to give **11** (488 mg, 92 % yield) as beige powder, mp = 51-52 °C; IR (KBr) ν 3093, 2957, 1728, 1567, 1416; ^1H NMR (CDCl_3) δ 3.93 (s 3H), 7.52 (d, 1H, J = 8.1 Hz), 7.93 (d, 1H, J = 8.1 Hz; ^{13}C NMR (CDCl_3) δ 53.5, 127.3, 128.7, 140.4, 141.7, 143.9, 165.0; Anal. Calcd for $\text{C}_7\text{H}_5\text{Br}_2\text{NO}_2$ (294.9): C, 28.51; H, 1.71; N, 4.75. Found: C, 29.01; H, 1.67; N, 4.71 %.

Procedure for nucleophilic substitution using sodium methoxide. To a stirred solution of methyl 2,6-dibromonicotinate (**11**, 1.5 g, 5.0 mmol) in dry MeOH (20 ml) was added NaOMe (270 mg, 5.0 mmol). The mixture was refluxed for 24 h and then poured into cold aq. NaHCO_3 (5 %, 50 ml) and the product was extracted with ether (3x20 mL). The separated organic phase was separated and concentrated *in vacuo*. Ether (40 mL) was added to the crude liquid and the organic phase was washed with brine (40 mL), dried (MgSO_4) and concentrated *in vacuo*. The

crude product was purified by chromatography on silica gel (CH_2Cl_2) to give a (7:3) mixture of **10** and **14** (923 mg, 75 %).

2-Bromo-6-methoxypyridine (12). To a stirred solution of 2,6-dibromopyridine (20 g, 84 mmol) in dry MeOH (50 mL) was added NaOMe (8g, 148 mmol). The mixture was refluxed for 25 h and then poured into a cold aq. soln. of NaHCO₃ (5%, 50 mL). The product was extracted with ether (3x30 ml) and the combined organic layers were washed with brine (40 ml) and concentrated *in vacuo*. The crude product was purified by chromatography on silica gel (EtOAc/Petrol 9:1) to give **12** (13 g, 83 %) as a liquid; bp=206-207°C; IR (KBr) ν 2953, 1596, 1582, 1558, 1472, 1413, 1298, 1022, 857; ¹H NMR (CDCl_3) δ 3.93 (s 3H), 6.69 (t, 1H, J = 7.7 Hz), 7.06 (d, 1H, J = 7.7 Hz), 7.40 (d, 1H, J = 7.7 Hz).

2-Bromo-6-methoxynicotinic acid (16). To a stirred solution of LDA (106 mmol) in dry THF (50 mL) was added dropwise under N₂ at -78°C a solution of 2-bromo-6-methoxypyridine **12** (10 g, 53 mmol) in dry THF (50 ml). After stirring 1 h. at the same temperature, the mixture was poured on an excess of carbonic dry ice. Solvents were removed *in vacuo* and the crude residue was dissolved in water (30 ml). The separated aqueous layer was washed with CH_2Cl_2 (3x15 ml) and pH was adjusted to 4 by adding aq. HCl (2M). The product was extracted with CH_2Cl_2 (3x15 ml) and the combined organic layers were washed with brine, dried (MgSO_4) and concentrated in *vacuo* to give **16** (1.6 g, 13 %) as a beige solid; mp= 212-213°C; IR (KBr) ν 1302, 1591, 1693, 2959, 3418; ¹H NMR (CDCl_3) δ 3.89 (s, 3H), 6.91 (d, 1H, J = 8.4 Hz), 8.10 (d, 1H, J = 8.4 Hz); ¹³C NMR (CDCl_3) δ 54.8, 109.4, 122.8, 137.9, 142.7, 163.7, 165.8. Anal. Calcd for C₇H₆BrNO₃ (232.0): C, 36.26; H, 2.61; N, 6.04. Found: C, 36.23; H, 2.67; N, 6.21 %.

Methyl 2-bromo-6-methoxynicotinate (10). To a stirred solution of 2-bromo-6-methoxynicotinic acid **16** (1 g, 4.3 mmol) and 3 drops of DMF in dry CH_2Cl_2 (20 ml) was slowly added oxalyl chloride (462 μ L, 5.4 mmol) at 0°C. The mixture was stirred at room temperature for 1 h and solvents were removed *in vacuo*. Dry methanol (30 ml) was then added at 0°C and the resulting mixture was stirred for 2 h at room temperature. Methanol was removed *in vacuo* and CH_2Cl_2 (30 ml) was added. The organic phase was washed with water (3x15 ml), dried (MgSO_4) and concentrated *in vacuo* to give the ester **10** (973 mg, 92 %) as white solid, mp= 54-55 °C; IR (KBr) ν 1247, 1586, 1724, 2952; ¹H NMR (CDCl_3) δ 3.85 (s, 3H), 3.93 (s, 3H), 6.93 (d, 1H, J = 8.5 Hz), 7.99 (d, 1H, J = 8.5 Hz); ¹³C NMR (CDCl_3) δ 42.5, 54.7, 109.3, 121.1, 139.7, 142.4, 164.5, 165.1; Anal. Calcd for C₈H₈BrNO₃ (246.0): C, 39.05; H, 3.28; N, 5.69. Found: C, 39.11; H, 3.32; N, 5.71 %.

Preparation of the cyclopenta[b]pyridin-2,5-dione (1)

(E)-Methyl 6-methoxy-2-(3-methoxy-3-oxoprop-1-enyl)nicotinate (9). A degassed mixture of methyl 2-bromo-6-methoxynicotinate (**10**, 0.30 g, 1.2 mmol), methyl acrylate (293 μ L, 3.3 mmol), allylpalladium chloride dimer Pd₂(allyl₂Cl₂) (24 mg, 0.065 mmol), P(o-Tol)₃ (40 mg, 0.13 mmol), Na₂CO₃ (320 mg, 3.0 mmol), toluene (2.53 ml) and dimethyl acetamide DMA (0.84 ml) was heated in a sealed tube at 115°C for 5 h. The reaction mixture was filtrated though Celite and concentrated in *vacuo*. The residue was purified by chromatography on silica gel (EtOAc /

Petrol 3:7) to give **9** (296 mg, 82%) as a yellow solid, mp= 112-113°C; IR (KBr) v 1132, 1243, 1482, 1589, 1737, 2961, 3012, 3097; ¹H NMR (CDCl₃) δ 3.81 (s, 3H), 3.90 (s, 3H), 3.99 (s, 3H), 6.72 (d, 1H, *J* = 8.6 Hz), 7.11 (d, 1H, *J* = 15.2 Hz), 8.11 (d, 1H, *J* = 8.1 Hz), 8.57 (d, 1H, *J* = 15.2 Hz); ¹³C NMR (CDCl₃) δ 52.2, 52.8, 54.2, 112.4, 119.7, 125.2, 140.9, 141.8, 152.8, 164.9, 166.5, 167.6; Anal. Calcd for C₁₂H₁₃NO₅ (251.2): C, 57.37; H, 5.22; N, 5.58. Found: C, 57.43; H, 5.34; N, 5.61 %.

Methyl 6-methoxy-2-(3-methoxy-3-oxopropyl)nicotinate (17). A degassed suspension of 10 % Pd/C (60 mg, 0.06 mmol) in a solution of **9** (150 mg, 0.6 mmol) in MeOH (10 ml) was vigorously stirred for 3 h at room temperature under H₂ (1 bar). The reaction mixture was filtered through a short pad of Celite and concentrated *in vacuo* to give **17** (141 mg, 93 %) as a yellow liquid; IR (KBr) v 1021, 1248, 1595, 1723, 2962; ¹H NMR (CDCl₃) δ 2.77 (t, 2H, *J* = 7.0 Hz), 3.47 (t, 2H, *J* = 7.0 Hz), 3.63 (s, 3H), 3.83 (s, 3H), 3.88 (s, 3H), 6.53 (d, 1H, *J* = 8.7 Hz), 8.06 (d, 1H, *J* = 8.7 Hz); ¹³C NMR (CDCl₃) δ 30.2, 30.7, 50.4, 52.6, 106.9, 116.6, 140.4, 159.9? 163.8, 165.5, 172.9; Anal. Calcd for C₁₂H₁₅NO₅ (253.2): C, 56.91; H, 5.97; N, 5.53. Found: C, 56.75; H, 6.04; N, 5.66 %.

Methyl 2-methoxy-5-oxo-6,7-dihydro-5*H*-cyclopenta[*b*]pyridine-6-carboxylate (18). Sodium methoxide (81 mg, 1.5 mmol) in THF (5 ml) was added to a solution of **17** (250 mg, 1 mmol) in dry THF (8 ml) under N₂ and the mixture was refluxed for 2 h. The pH was then adjusted to 5 by adding aq. HCl (2M) before extraction of the product with CH₂Cl₂ (2x15 ml). The combined organic layers was washed with aq. Na₂CO₃ (10%, 5 ml), sat. aq. NH₄Cl (5 ml), dried (MgSO₄) and concentrated *in vacuo*. The crude product was purified by chromatography on silica gel (EtOAc / petrol 3:7) to give **18** (166 mg, 75 %) as a beige solid, mp= 81-82 °C; IR (KBr) v 1021, 1250, 1439, 1595, 1736, 2960; ¹H NMR (CDCl₃) δ 3.29-3.31 (m, 1H), 3.45-3.50 (m, 1H), 3.70-3.74 (m, 1H), 3.73 (s, 3H), 4.00 (s, 3H), 6.68 (d, 1H, *J* = 8.7 Hz), 7.82 (d, 1H, *J* = 8.7 Hz); ¹³C NMR (CDCl₃) δ 33.3, 53.1, 53.2, 54.9, 112.2, 123.3, 135.1, 169.7, 169.8, 175.1, 196.5.

6,7-Dihydro-1*H*-cyclopenta[*b*]pyridine-2,5-dione (1). A solution of **18** (100 mg, 0.45 mmol) in aq. HCl (5 M, 2 ml) was refluxed for 3 h. The mixture was cooled to room temperature and the pH was adjusted to 5 by adding aq. K₂CO₃ (2 M). The product was extracted with EtOAc (5x15mL). The combined organic layers were dried (MgSO₄), concentrated *in vacuo* to give **1** (56 mg, 84 %) as a beige solid, mp> 260°C; IR (KBr) v 3085, 2924, 1675-1653, 1427, 1105; ¹H NMR (DMSO) δ 2.56 (m, 2H), 2.94 (t, 2H, *J* = 5.1 Hz), 6.31 (d, 1H, *J* = 9.4 Hz), 7.58 (d, 1H, *J* = 9.4 Hz), 12.6 (s, 1H); ¹³C NMR (DMSO) δ 24.5, 34.7, 116.3, 119.2, 134.1, 164.1, 169.5, 199.3; Anal. Calcd for C₈H₇NO₂ (149.1): C, 64.43; H, 4.73; N, 9.39. Found: C, 64.42; H, 4.83; N, 9.23 %.

References and Notes

1. Fossa, P.; Menozzi, G.; Dorigo, P.; Floreani, M.; Mosti, L. *Bioorg. Med. Chem.* **2003**, *11*, 4749. The authors reported that cyclopenta[b]pyridin-2,5-dione is able to increase atrial contractility to an extent which is significantly higher than of Milrinone. Cyclopenta[b]pyridin-2,5-dione is similar to Milrinone in its pharmacological action and in its activity. The positive inotropic effect exerted is related to an increase of cAMP levels obtained through inhibition of cGMP-inhibited cAMP phosphodiesterase (PDE) called Type 3 (PDE3). The maximal inhibition of cyclopenta[b]pyridin-2,5-dione against PDE3 being $81.93 \pm 0.29\%$ versus 100% of Milrinone.
2. Prakash, H. S.; Senthilkumar, S. P. *Curr. Org. Chem.* **2004**, *8*, 1521.
3. Ramirez, F.; Paul, A. P. *J. Org. Chem.* **1954**, *19*, 183.
4. Robert, N.; Hoarau, C.; Célanire, S.; Ribéreau, P. ; Godard, A. ; Guéguiner, G. ; Marsais, F. *Tetrahedron* **2005**, *61*, 4569.
5. Hirokawa and co-workers (Hirokawa, Y.; Horijawa, T.; Kato, S. *Chem. Pharm. Bull.* **2000**, *48*, 1847-1853 previously reported that the regioselective displacement of a chlorine from methyl 2,6-dichloronicotinate with sodium methoxide is more effective using MeOH as solvent leading to methyl 2-chloro-6-methoxychloroniconinate as the major isomer (7:3 ratio).
6. Above -50°C the C-3 lithio- anion was degraded.
7. Mutterer, F.; Weis, C.; Claus, D. *Helv. Chim. Acta* **1976**, *59*, 229.