Synthesis and structural characterisation of amides from picolinic acid and pyridine-2,6-dicarboxylic acid

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Coupling picolinic acid (pyridine-2-carboxylic acid) and pyridine-2,6-dicarboxylic acid with N-alkylanilines affords a range of mono- and bis-amides in good to moderate yields. These amides are of interest for potential applications in catalysis, coordination chemistry and molecular devices. The reaction of picolinic acid with thionyl chloride to generate the acid chloride in situ leads not only to the N-alkyl-N-phenylpicolinamides as expected but also the corresponding 4-chloro-N-alkyl-N-phenylpicolinamides in the one pot. The two products are readily separated by column chromatography. Chlorinated products are not observed from the corresponding reactions of pyridine-2,6-dicarboxylic acid. X-Ray crystal structures for six of these compounds are described. These structures reveal a general preference for cis amide geometry in which the aromatic groups (N-phenyl and pyridyl) are cis to each other and the pyridine nitrogen anti to the carbonyl oxygen. Variable temperature 1H NMR experiments provide a window on amide bond isomerisation in solution.

Results & Discussion

Synthesis. Amides 5a–c, 6a–c, 7a–c, 8a–b were prepared by activating picolinic acid 3 and pyridine-2,6-dicarboxylic acid 4 to the corresponding acid chlorides in situ, or via diimide-mediated peptide coupling. Activating picolinic acid 3 with thionyl chloride afforded not only the simple picolinamides 5a–c as expected, but also the 4-chloropicolinamides 6a–c in the same pot. The two products were easily separated by column chromatography, enabling a ‘two for the price of one’ synthesis of new amides.

The mono-amide ligands 5a–c and 6a–c were synthesised from picolinic acid 3 and the corresponding aniline in one pot, via the acid chloride (Figure 3). Thus acid 3 was treated with thionyl chloride overnight, followed by N-methylaniline, N-ethylaniline or N-diphenylamine and triethylamine in dichloromethane. This route gave the anticipated products 5a–c in low to moderate yields (31–54%), and the 4-chloro derivatives 6a–c, isolated in small but utilisable yields (10–13%). Each pair of compounds was readily separated by column chromatography.

Chlorination of the ring presumably occurs via activation of the pyridine to nucleophilic attack by chloride anion. This could occur during formation of the acid chloride or in the subsequent coupling step. The direct synthesis of 4-chloropicolinyl chloride from picolinic acid using thionyl chloride has been reported previously, although in our own prior work we have converted picolinic acid to picolinoyl chloride with this reagent system, then reacted the acyl chloride with L-proline, without observing ring-chlorinated side products.
Our efforts to characterise the acid chloride intermediate(s) were unsuccessful: we were able to isolate a low-melting orange solid (mp 40–50 °C) but this quickly decomposed before it could be further characterised.

The N-methyl mono-amide 5a has been prepared previously by Habib and Rees, who reported its synthesis, melting point and elemental analysis\(^{26}\), and more recently by Okamoto et al. as part of an investigation into acid-induced conformational changes in aromatic amides\(^ {14}\). Habib and Rees prepared 5a via the acid chloride, reacting picolinic acid 3 and thionyl chloride in benzene, then adding N-methylaniline dropwise and heating at reflux; Okamoto activated acid 3 as the mixed anhydride by reaction with ethyl chloroformate and triethylamine, before adding N-methylaniline. The 4-chloro derivative 6a was not isolated in either of these previous syntheses.

Bis-amides 7a–c were prepared in a similar manner, from pyridine-2,6-dicarboxylic acid 4 in one pot (Figure 3b). This gave compounds 7a–c as crystalline solids in excellent yield (86–90%); chlorinated byproducts were not observed from the reactions of dicarboxylic acid 4. Compounds 7a and 7b appear previously in the literature, but details of their synthesis and characterisation are incomplete. Ried and Neidhardt studied “hydrogenolysis” of the N-methyl compound 7a and related quinoline carboxylic acids upon reaction with lithium aluminium hydride\(^ {27}\). The N-methyl (7a) and N-ethyl (7b) analogues have been used to generate metal complexes\(^ {17,18}\) and in metal extraction experiments\(^ {19–21}\), while Dobler et al. conducted computational experiments to describe the interaction between ligands of this type and lanthanide cations\(^ {28}\). Kapoor and coworkers recently reported synthesis and structural characterisation of related thioamide derivatives\(^ {29}\).

In a complementary approach, the peptide derivatives 8a and 8b were prepared from pyridine-2,6-dicarboxylic acid 4 using diimide coupling methodology\(^ {23}\). Thus dipeptides L-valinyl-S-benzyl-L-cysteine methyl ester (tosylate salt) 9 and S-benzyl-L-cysteinyl-L-valine methyl ester 10 (prepared from L-cysteine and L-valine via

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**Figure 1** | General structures of bidentate 1 and tridentate 2 amide targets, prepared from picolinic acid 3 and pyridine-2,6-dicarboxylic acid 4.

**Figure 2** | Structures of target amides 5–8. For 5a–7a R = Me, 5b–7b R = Et, 5c–7c R = Ph; 8a is derived from the l-valinyl-l-cysteine dipeptide (R’ = ’Pr, R’ = CH₂SBn), while 8b incorporates the l-cysteinyl-l-valine dipeptide (R’ = CH₂SBn, R’ = ’Pr).

**Figure 3** | Synthesis of target compounds. (a) Synthesis of mono-amides 5a–c and 6a–c: i. SOCl₂, reflux, 16 h; ii. Et₃N, N-methylaniline a; N-ethylaniline b or N-diphenylamine c, DCM, rt, 16 h; 5a 35%/6a 13%; 5b 31%/6b 10%; 5c 54%/6c 10% (yields over two steps for major/minor products). (b) Synthesis of bis-amides 7a–c: i. SOCl₂, reflux, 16 h; ii. Et₃N, N-methylaniline a, N-ethylaniline b or N-diphenylamine c (2 eq.), DCM, rt, 16 h; 7a 86%, 7b 88%, 7c 90% (over two steps). (c) Synthesis of peptide derivatives 8a–b: iii. EDCI, HOBt, Et₃N, L-valinyl-S-benzyl-l-cysteine methyl ester tosylate salt 9 or S-benzyl-l-cysteinyl-l-valine methyl ester 10 (prepared from L-cysteine and l-valine via
standard methods\(^{16}\)) were coupled with 4 to give the peptide derivatives 8a and 8b in moderate yields (Figure 3c).

**Crystallographic investigations.** The geometry of the amide bond in compounds such as these has received attention previously with a view to potential applications in molecular switches and devices\(^{14–16}\). N-Alkylation – specifically N-methylation – has been shown to induce a change from trans-preferential to cis-preferential amides (Figure 4).

Thus while the amide bond in benzanilide 11 (R = H) is trans, the corresponding bond in N-methylenzanilide 12 (R = Me) is preferentially cis, both in the crystalline state and in solution\(^{14}\). Likewise crystallographic and NMR characterisation of 5a reported by Okamoto et al. show that the two aromatic groups adopt a cis relationship in that compound too\(^{16}\). To investigate the geometry of the amides prepared in the current study, single crystal X-ray structures were determined for the mono-amides 5b and 5c, 4-chloro mono-amides 6b and 6c, and bisamides 7a and 7c (Figures 5 and 6; Supplementary Information).

The structures of the N-methyl (7a) and N-ethyl (5b, 6b) compounds reveal cis amide geometry in all cases: the aromatic groups (N-phenyl and pyridyl) are cis to each other, and the methyl or ethyl substituent is cis to the carbonyl group. There is also a general preference for the pyridine nitrogen to sit anti to the carbonyl oxygen(s). Among the mono-amides, these groups are antical in 5b (the O–C–C–N dihedral angle is 123.9\(^\circ\)), 6b (126.5\(^\circ\)) and 6c (137.6\(^\circ\)), but synclinal in 5c (56.7\(^\circ\)) (Figure 5). Of the bis-amide structures, the pyridine nitrogen is antical to both carbonyls in the tetraphenyl compound 7c; there are two inequivalent molecules of 7c in the crystal structure, which exhibit dihedral angles around the bond in question of 141.6\(^\circ\) and 131.9\(^\circ\)/139.1\(^\circ\) and 149.8\(^\circ\) respectively. However in the dimethyl compound 7a, the pyridine nitrogen is anti to one of the amide carbonyls (137.2\(^\circ\)) but syn to the other (−57.2\(^\circ\)), which – in combination with the two cis amide bonds – positions the two phenyl groups in close proximity and an edge-to-face arrangement (Figure 6).

**Variable temperature NMR experiments.** In light of the recent work by Okamoto et al. using \(^1\)H NMR to follow cis/trans isomerisation in related aromatic amides\(^{14}\), we were interested to note evidence for slow conformational change in the \(^1\)H NMR spectra of compounds 7a–c. The room temperature \(^1\)H NMR spectra of 7a–c are generally poorly resolved with considerable line broadening (in contrast to the spectra of corresponding monoamides 5a–c in which equivalent line broadening is not observed – see Supplementary Information). Variable temperature \(^1\)H NMR data for the ethyl substituted ligand 7b (Figure 7) shows that signals resolve as the temperature is increased, confirming that the observed line broadening arises due to slow conversion between amide conformational isomers at room temperature. For example the signal at ~ 3.7 ppm, due to the methylene protons of the ethyl group, is a broad apparent singlet at 300 K but a clearly resolved quartet at 350 K (see inset in Figure 7).

**Conclusion.** Amides derived from picolinic acid 3 and pyridine-2,6 dicarboxylic acid 4 have potential applications in catalysis, coordination chemistry and molecular switches. These compounds are readily prepared via the acid chloride or applying peptide coupling reagents. X-Ray crystal structures reveal that the generally preferred geometry of these amides positions the aromatic groups cis to each other and the pyridine nitrogen anti to the carbonyl oxygen. Variable temperature NMR experiments indicate slow cis/trans isomerisation in solution for the bis-amide series.

**Methods**

**Amide synthesis.** General procedure 1. Thionyl chloride (8.0 mL, 109 2 mmol) was added to picolinic acid 3 (1.00 g, 8.20 mmol) and the resulting suspension was refluxed for 16 h. The orange coloured solution was reduced in vacuo to give the acid chloride as a bright orange oil. The oil was dissolved in dry DCM (40 mL) and cooled to 0 °C. A solution of N-alkylaniline (16.20 mmol) and triethylamine (2.20 mL, 16.20 mmol) in dry DCM (20 mL) was added via cannula. The resulting purple coloured solution was stirred at 0°C for 20 min and at room temperature for 16 h after which time the solution had become dark brown. The solution was washed with half-saturated aqueous ammonium chloride solution (2 × 12 mL), water (2 × 6 mL) and dried (Na2SO4), then concentrated in vacuo.

**General procedure 2.** Thionyl chloride (4.0 mL, 66 mmol) was added to 2,6-pyridinedicarboxylic acid 4 (0.50 g, 3.0 mmol) and the resulting suspension was refluxed under an argon atmosphere for 16 h to give a clear yellow oil. Excess thionyl chloride was removed in vacuo and the acid chloride was dissolved in dry CH2Cl2 (10 mL) and cooled to 0 °C. A solution of N-alkylaniline (12.0 mmol) and triethylamine (0.84 mL, 6.0 mmol) in dry DCM (2.5 mL) was added via cannula. The resulting mixture was stirred at room temperature for 16 h during which time a white precipitate formed. The suspension was washed with half-saturated aqueous ammonium chloride solution (2 × 6 mL) and water (2 × 3 mL), then dried (Na2SO4) and concentrated in vacuo.

**General procedure 3.** Pyridine-2,6-dicarboxylic acid 4 (0.10–0.30 g, 1 eq.), dipetide amine (as the free amine or tosylate salt, 2 eq.), 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDCI, 2 eq.) and 1-hydroxybenzotriazole (HOBT, 2 eq.) were dissolved in DCM (10–30 mL) and triethylamine (2 eq. for free amine, 4 eq. for tosylate salt) was added. The reaction mixture was stirred at room temperature for 22–48 h while monitored by TLC. Additional DCM or chloroform (10–20 mL) was added and the solution washed with equivalent volumes of water, 1 M hydrochloric acid, saturated sodium bicarbonate (aqueous) and brine (4 M MgSO4) then concentrated in vacuo.
N-Methyl-N-phenylpicolinamide \(5a\) and 4-Chloro-N-methyl-N-phenylpicolinamide \(6a\). Picolinic acid \(3\) (1.0 g, 8.2 mmol) and N-methylaniline (1.76 mL, 16.2 mmol) were coupled using thionyl chloride (Procedure 1). TLC of the crude mixture showed the presence of two products, which were separated by flash column chromatography (petroleum benzine/ethyl acetate, 1:1) to afford \(5a\) (0.60 g, 35%) as a white crystalline solid and \(6a\) (0.27 g, 13%) as a thick, clear, colourless oil.

Data for N-methyl-N-phenylpicolinamide \(5a\) in agreement with literature\(^1\); see Supplementary Information for details.

Figure 6 | Crystal structures of amides 7a (CCDC-1002450) and 7c (1002451). Carbon atoms are shown in grey, oxygen in red, nitrogen in blue and hydrogen in white. In 7a the methyl group and carbonyl oxygen are cis. The pyridine nitrogen is anti to both carbonyl oxygen atoms in 7c, but syn to one and anti to the other in 7a.

Figure 7 | Variable temperature \(^1\)H NMR spectra of bis-amide ligand 7b (400 MHz, d\(_8\)-toluene), confirming slow conformational change at room temperature.
Data for 4-chloro-N-methyl-N-phenylpicolinamide 6a. R = 0.04 (petroleum benzene/ethyl acetate, 1:1); mp = 123–129 °C (C8H17CONH-CH2CO2H, cm−1) 3058 (w), 1670 (s), 1587 (m), 1488 (m); δH 0.0 (CH2CO2H) 7.19–7.32 (10H, m, C(C3H2)2), 7.63 (H, d, J = 1.0 Hz, 1× pyr-CH), 7.77 (H, t, J = 7.5 Hz, 1× pyr-CH), 8.28–8.30 (2H, m, pyr-CH); δC (100 MHz, CDCl3) 123.9, 124.7, 126.9, 128.9, 134.3, 134.7, 146.9, 156.9, 166.9; m/z (ES+) 275 (100%, [M+Na]+), 297 (55%, [M]+); HRMS (ES+) C16H13N2O3Na+; [M + Na]+ requires 297.09984, found 297.09985.

Data for 4-chloro-N-methyl-N-phenylpicolinamide 6c. R = 0.03 (petroleum benzene/ethyl acetate, 1:1); mp = 123–129 °C (C8H17CONH-CH2CO2H, cm−1) 3058 (w), 1670 (s), 1587 (m), 1488 (m); δH 0.0 (CH2CO2H) 7.19–7.32 (10H, m, C(C3H2)2), 7.63 (H, d, J = 1.0 Hz, 1× pyr-CH), 7.77 (H, t, J = 7.5 Hz, 1× pyr-CH), 8.28–8.30 (2H, m, pyr-CH); δC (100 MHz, CDCl3) 123.9, 124.7, 126.9, 128.9, 134.3, 134.7, 146.9, 156.9, 166.9; m/z (ES+) 275 (100%, [M+Na]+), 297 (55%, [M]+); HRMS (ES+) C16H13N2O3Na+; [M + Na]+ requires 297.09984, found 297.09985.

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N,N-Diphenylpyridazin-5-yl acetate, 5. Barry, S. M. & Rutledge, P. J. cis-dihydroxylation of alkenes by a non-heme iron donors: Synthesis, structure, and reactivities. J. Am. Chem. Soc. 123, 882–887 (2001).

Pyrindine-2,6-dicarboxylic acid bis(2-phenyl-cysteine methyl ester)carbazole 8a. Pyridinetricarboxylic acid 4 (0.10 g, 0.58 mmol) and 2-phenyl-cysteine methyl ester tosylate salt 9 (0.60 g, 1.2 mmol) were coupled using EDCI/HOBt (Procedure 3) to give 8a as an yellow oil (0.2 g, 44%) after purification by column chromatography (cyclohexane/ethyl acetate, 1:1); R = 0.65 (cyclohexane/ethyl acetate, 1:1); [α]20D = +15 (c = 0.2, CH2Cl2), C6H5CONH2 (thin film) 3290, 1745, 1659 (s), 1236 (w), 1216 (12H, d, J = 6.5 Hz, 2× CH2Cl2), 2.25–2.37 (2H, m, 2× CH2Cl2), 2.86–2.88 (4H, m, 2× CH2SCPh2), 3.67 (H, s, 2× CH2), 7.29 (4H, d, J = 7.3 Hz, 2× CH2), 7.92 (1H, m, pyr-CH); δC (100 MHz, CDCl3) 125.4, 127.1, 128.5, 137.9, 144.7, 153.8, 167.9; m/z (ES+) 470 (100%, [M]+), 492 (43%, [M]+); HRMS (ES+) C30H29N2O9; [M]+ requires 470.18631, found 470.18615.

Pyrindine-2,6-dicarboxylic acid bis(2-phenyl-cysteine methyl ester)carbazole 8b. Pyridinetricarboxylic acid 4 (0.25 g, 1.5 mmol) and 2-phenyl-cysteine methyl ester tosylate salt 9 (1.0 g, 3.0 mmol) were coupled using EDCI/HOBt (Procedure 3) to give 8b as an yellow oil (1.13 g, 61%), after purification by column chromatography (cyclohexane/ethyl acetate, 1:1); R = 0.55 (cyclohexane/ethyl acetate, 1:1); [α]20D = −7.6 (c = 0.2, CH2Cl2), C6H5CONH2 (thin film) 3420, 2390 (s, br), 2390 (s), 1740 (s), 1629 (w), 1271 (2H, d, J = 4.5 Hz, 2× CH2Cl2), 2.11–2.22 (2H, m, 2× CH2Cl2), 2.91 (H, dd, J = 14.0, 7.5 Hz, 2× CH2Cl2), 3.74 (H, s, 2× OCH3), 3.84 (H, t, 2× CH2), 4.51 (2H, dd, J = 8.5, 5.0 Hz, 2× CH2Cl2), 4.69–4.77 (2H, m, 2× CH2Cl2), 6.94 (2H, d, J = 8.5 Hz, 2× CH2), 7.17–7.70 (10H, m, C6H5), 8.02–8.10 (2H, m, pyr-CH), δC (100 MHz, CDCl3) 125.4, 127.1, 128.5, 137.9, 144.7, 153.8, 167.9; m/z (ES+) 780 (100%, [M]+); HRMS (ES+) C35H27N3O9S5; [M]+ requires 780.3101, found 780.3112.
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**Author Contributions**

S.M.B., K.M.H. and P.J.R. conceived and designed the experiments. P.D., S.M.B. and K.M.H. performed the synthetic experiments; M.J.M., P.T. and P.J. conducted X-ray crystallography experiments. M.I.M., P.T. and P.J. (crystallography), P.D., S.M.B., K.M.H. and P.J.R. analyzed the data. S.M.B. and P.J.R. wrote the main manuscript text including Figures 1–4 and 7; P.D. and M.J.M. prepared figures 5 and 6. All authors reviewed the manuscript.

**Additional information**

Accession codes: Crystallographic data are contained in CCDC-1002446 (5b), -1002447 (5c), -1002448 (6b), -1002449 (6c), -1002450 (7a) and -1002451 (7c). These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

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