

[W&M ScholarWorks](https://scholarworks.wm.edu/)

[Arts & Sciences Articles](https://scholarworks.wm.edu/aspubs) **Articles** [Arts and Sciences](https://scholarworks.wm.edu/as) Articles Arts and Sciences Arts and Sciences Articles **Arts** and Sciences Arts and Sciences **Arts** and Sciences **Arts** and Sciences **Arts** and Sciences **Arts** and Scien

2008

Poly[l-2-aminopyrazine-j2 N1 :N4 - l-cyanido-copper(I)]: A Threedimensional Network From Laboratory Powder Diffraction Data

Silvina Pagola

Robert D. Pike William & Mary, rdpike@wm.edu

Kathryn deKrafft

Tristan A. Tronic

Follow this and additional works at: [https://scholarworks.wm.edu/aspubs](https://scholarworks.wm.edu/aspubs?utm_source=scholarworks.wm.edu%2Faspubs%2F1978&utm_medium=PDF&utm_campaign=PDFCoverPages)

Part of the Chemistry Commons

Recommended Citation

Pagola, Silvina; Pike, Robert D.; deKrafft, Kathryn; and Tronic, Tristan A., Poly[l-2-aminopyrazine-j2 N1 :N4 l-cyanido-copper(I)]: A Three-dimensional Network From Laboratory Powder Diffraction Data (2008). Acta Crystallographica Section C, C64. https://doi.org/10.1107/S010827010800231X

This Article is brought to you for free and open access by the Arts and Sciences at W&M ScholarWorks. It has been accepted for inclusion in Arts & Sciences Articles by an authorized administrator of W&M ScholarWorks. For more information, please contact scholarworks@wm.edu.

metal-organic compounds

Acta Crystallographica Section C Crystal Structure Communications ISSN 0108-2701

Poly[μ -2-aminopyrazine- $\kappa^2 N^1$: N^4 - μ -cyanido-copper(I)]: a three-dimensional network from laboratory powder diffraction data

Silvina Pagola,^a* Robert D. Pike,^b Kathryn deKrafft^b and Tristan A. Tronic^b

^aCollege of William and Mary, Physics Department, Williamsburg, VA 23187, USA, and Applied Research Center, 12050 Jefferson Avenue, Newport News, VA 23606, USA, and ^bCollege of William and Mary, Chemistry Department, Williamsburg, VA 23187, USA

Correspondence e-mail: spagol@wm.edu

Received 14 September 2007 Accepted 22 January 2008 Online 16 February 2008

In the title compound, $\left[\text{Cu(CN)}(\text{C}_{4}\text{H}_{5}\text{N}_{3})\right]_{n}$ or $\left[\text{Cu}(\mu\text{-CN)}(\mu\text{-}1)\right]_{n}$ $PyzNH₂$)]_n (PyzNH₂ is 2-aminopyrazine), the Cu^I center is tetrahedrally coordinated by two cyanide and two PyzNH2 ligands. The Cu^I-cyano links give rise to [Cu–CN]_{∞} chains running along the c axis, which are bridged by bidentate PyzNH₂ ligands. The three-dimensional framework can be described as being formed by two interpenetrated threedimensional honeycomb-like networks, both made of 26 membered rings of composition $\left[\text{Cu}_{6}(\mu\text{-CN})_{2}(\mu\text{-PyzNH}_{2})_{4}\right]$.

Comment

Metal-organic networks of Cu^I-cyano and bridging diimine ligands have potential applications in gas storage and catalysis. Metal–organic materials containing luminescent metals, such as copper(I), and having suitable network porosity can potentially be used in gas molecule sensing systems, since the inclusion of small molecules into the network can alter the luminescent behavior of the material. The wide variety of bonding modes of Cu^I-cyano units allows the preparation of new materials of a large number of structural types, where the Cu^I coordination often varies from 2 to 5. Furthermore, the assembly of Cu^I and asymmetrically substituted diimine ligands can lead to chiral two- and three-dimensional networks, which could find applications in asymmetric catalysis and nonlinear optics (Teichert & Sheldrick, 1999, and references therein).

We have recently prepared a variety of new Cu^I-cyano diimine compounds by open reflux reactions and hydrothermal syntheses, and have investigated their luminescence properties (Tronic et al., 2007). The ligands studied include pyrazine (Pyz), 2-aminopyrazine (PyzNH₂), quinoxaline, phenazine, 4,4'-bipyridyl, pyrimidine, 2-aminopyrimidine, 2,4diaminopyrimidine, 2,4,6-triaminopyrimidine, quinazoline, pyridazine and phthalazine. As part of this study, poly $\lceil u^2 - 2 \rceil$ aminopyrazine- μ -cyanido-copper(I)], $[CuCN(\mu-PyzNH_2)]_n$, (I), was prepared by an open reflux reaction. This reaction did not produce single crystals suitable for structure determination, but the structure of (I) has been solved from the X-ray powder diffraction pattern collected at room temperature.

(1) Only a few 1:1 $\left[Cu(CN)(\mu\text{-ligand}) \right]$ complexes have been previously reported, all containing Cu^I with distorted tetrahedral coordination. In the crystal structure of $\left[\text{CuCN}(\mu - \text{Hilb}) \right]$ $p \cdot (p \cdot d \cdot b)$ is di-4-pyridyl-*trans*-1,4-divinylbenzene), the long bidentate pdvb ligand favors the formation of corrugated sheets, each sheet being composed of parallel $\left[\text{Cu–pdvb}\right]_{\infty}$ zigzag chains (Hanika-Heidl et al., 2003). The sheets are linked together through cyano bridges that give rise to [Cu–CN]_{∞}

Figure 1

(a) $\text{[Cu}_6(\text{CN})_2(\mu\text{-Pyz})_4$] 26-membered rings in the three-dimensional honeycomb-like arrays of $[Cu(CN)(\mu-Pyz)]_n$. H atoms are not shown for clarity. (b) A view of the distorted tetrahedral coordination around the Cu^I atom in $[Cu(CN)(\mu$ -PyzNH₂)]_n, showing the atom-labeling scheme (H-atom labels have been omitted for clarity).

Figure 2

The final Rietveld refinement. Observed intensity (points), calculated profile (solid line) and peak positions (| symbols). The difference plot $(I_{observed} - I_{calculated})$ is shown at the bottom.

chains running along the direction perpendicular to the plane of the sheets. This structure exhibits remarkably different $Cu^I \cdot ^{\alpha}$ Cu^I separations of 4.804 (via CN⁻) and 20.366 Å (via pdvb). The pdvb Cu—N interatomic distances are 2.249 and 2.173 Å, whereas the cyano $Cu-C$ and $Cu-N$ distances are 1.880 and 1.941 \AA , respectively.

 $[Cu(CN)(\mu$ -2-MePyz)]_n (2-MePyz is 2-methylpyrazine) and $[Cu(CN)(\mu$ -4-MePym)]_n (4-MePym is 4-methylpyrimidine) form three-dimensional frameworks in which one-dimensional $\left[\text{Cu}-\text{CN}\right]_{\infty}$ chains are bridged by the linear bidentate aromatic ligands (Teichert & Sheldrick, 1999). In $\left[\text{Cu(CN)}\right]\mu$ -2-MePyz)]_n, 22-membered rings of $[Cu₆(CN)₄(\mu$ -2-MePyz)₂] composition form a chiral honeycomb-like two-dimensional network; these units are connected by additional 2-MePyz ligands, giving rise to a porous and chiral three-dimensional framework in $P2_12_12_1$. On the other hand, the linear ligand 4-MePym bonds to Cu^I atoms at shorter distances owing to its intramolecular geometrical disposition of N atoms, and it forms tetramers in a centrosymmetric structure containing disordered cyano C and N atoms. Furthermore, regardless of the asymmetric nature of the 4-MePym ligand, $\left[\text{Cu(CN)}\right]\mu$ -4-MePym)]_n crystallizes in the nonchiral space group $P4₂/n$.

 $[Cu(CN)(\mu-Pyz)]_n$ forms a three-dimensional network in the $P2_1/c$ space group [a = 6.208 (2) Å, b = 9.158 (2) Å, c = 11.198 (2) \AA and $\beta = 90.89$ (3)°], wherein each Cu^I center is tetrahedrally coordinated to two cyano and two Pyz units. [Cu–CN]_{∞} chains run along the b axis and are connected by two Pyz ligands, giving rise to 22-membered rings of composition $\left[\text{Cu}_6(\text{CN})_4(\mu\text{-Pyz})_2\right]$. Alternatively, we can describe the three-dimensional network as formed by two interpenetrated, but otherwise symmetrically equivalent, three-dimensional honeycomb-like arrays made of 26-membered rings of composition $\left[\text{Cu}_6(\text{CN})_2(\mu\text{-Pyz})_4\right]$ (Fig. 1*a*). Offset face-to-face $\pi-\pi$ interactions within Pyz ligands are present, with a

centroid-to-centroid distance of 4.58 (6) \AA . The Pyz Cu^I $-N$ distances are $2.159(12)$ and $2.134(12)$ Å, whereas the cyano Cu^I – C and Cu^I – N distances are 1.920 (18) and 1.944 (19) \AA , respectively, and the Cu^I tetrahedral angles are 99.3 (5). 133.1 (8), 99.8 (6), 111.1 (8), 101.4 (6) and 107.2 (8)°. The existence of some C/N disorder (not refined) in the cyano ligands is reported for this structure (Kuhlman et al., 1999).

A similar packing was found for (I). The refined unit-cell parameters (after axes permutations) are also similar, even though $\left[\text{Cu(CN)}(\mu\text{-Pyz})\right]_n$ and $\left[\text{Cu(CN)}(\mu\text{-PyzNH}_2)\right]_n$ belong to the monoclinic and orthorhombic crystal systems, respectively. Fig. 1(b) shows the distorted tetrahedral coordination adopted by Cu1. This figure also shows one of the two interpenetrated three-dimensional honeycomb-like arrays formed by 26-membered rings of $\left[\text{Cu}_6(\text{CN})_2(\mu\text{-PyzNH}_2)_4\right]$ composition, which in turn generate the three-dimensional network of $[Cu(CN)(\mu-PyzNH_2)]_n$. The compound crystallizes in the noncentrosymmetric and nonchiral space group $Pca2₁$ (No. 29), and the cyano C and N positions for (I) refine as ordered. Additionally, offset face-to-face $\pi-\pi$ interactions between the PyzNH2 ligands with a centroid-to-centroid distance of 4.16 (5) Å are found.

In view of the chemical similarity of Pyz and PyzNH2, and the similarity of the crystal packings of $[CuCN(\mu-Pyz)]_n$ and $[CuCN(\mu-PyzNH_2)]_n$, it is reasonable that the ligand Cu–N distances found are very close for the two complexes. The differences in the tetrahedral coordination angles around $Cu¹$ are slightly larger.

Experimental

Compound (I) was prepared by open reflux synthesis as described by Tronic et al. (2007). Equimolar quantities of copper(I) cyanide and KCN were suspended in water and warmed. PyzNH₂ was added in half of the previous molar quantity and the suspension was refluxed overnight under a nitrogen atmosphere. The reaction mixture was filtered, and the solid obtained was washed with water, ethanol and diethyl ether and dried under vacuum. A yellow powder was isolated. The C, H and N elemental compositions were measured by standard techniques, and the Cu content was determined by atomic absorption spectroscopy.

Data collection

Philips Analytical X'Pert Pro MRD diffractometer

Specimen mounting: packed powder in flat plate holder

Specimen mounted in reflection mode

Scan method: step $2\theta_{\min} = 14.0, 2\theta_{\max} = 135.0^{\circ}$ Increment in $2\theta = 0.03^{\circ}$

metal-organic compounds

Refinement

The powder diffraction pattern was indexed without impurity peaks with the program DICVOL04 (Boultif & Louër, 2004). Le Bail fits (Le Bail et al., 1988) performed with the program GSAS (Larson & Von Dreele, 2000) confirmed the validity of the orthorhombic unitcell parameters. Starting with coordinates for Cu1, the program PSSP (Powder Structure Solution Program; P. W. Stephens & S. Pagola; http://powder.physics.sunysb.edu/programPSSP/pssp.html), using direct-space methods and the simulated annealing algorithm, was applied to determine the location of the cyano and $PyzNH₂$ ligands. The atomic positions of the $PyzNH₂$ ligand were refined as a rigid body [geometry from Cambridge Structural Database (Allen, 2002) entry AMPYRZ (Chao et al., 1976)], and H-atom positions were calculated with the program $WinGX$ (Farrugia, 1999). A positivedefinite set of anisotropic displacement parameters could be determined for the Cu1 atom, whereas the values of the remaining non-Hatom isotropic displacement parameters were refined subject to a group constraint. $U_{\text{iso}}(H)$ values were constrained to a value of 1.2 times the equivalent isotropic displacement parameter of the attached non-H atom.

A bond-length restraint of 1.160 (1) \AA was included in order to obtain a reasonable cyano bond length, although no restraints were necessary for PyzNH2 Cu—N distances or angles. The possibility of disorder in the cyano C9 and N10 atoms was investigated by switching C- and N-atom identities. It was concluded that the cyano C and N positions are ordered.

The following parameters were constrained to equivalent values for the three histograms: lattice parameters, 2θ zero error, transparency, sample displacement error, profile parameters, atomic positions, isotropic and Cu1 anisotropic displacement parameters, preferred orientation and absorption coefficients. The standard deviations of the crystallographic parameters have been corrected following the procedure reported by Scott (1983). A plot of the observed and calculated powder diffraction intensities and their difference (at the bottom) is shown in Fig. 2.

Data collection: *X'Pert Data Collector* (Philips, 2002); cell refinement: GSAS (Larson & Von Dreele, 2000); data reduction: GSAS; program(s) used to solve structure: EXPO2004 (Altomare et al., 2004) and PSSP (http://powder.physics.sunysb.edu/programPSSP/ pssp.html); program(s) used to refine structure: GSAS; molecular graphics: Mercury (Macrae et al., 2006) and ORTEP-3 (Farrugia, 1997); software used to prepare material for publication: publCIF (Westrip, 2008).

SP thanks to the Department of Physics of the College of William and Mary for funding. RDP gratefully acknowledges the donors of the American Chemical Society Petroleum Research Fund (grant No. 44891-B3).

Supplementary data for this paper are available from the IUCr electronic archives (Reference: FA3122). Services for accessing these data are described at the back of the journal.

References

- Allen, F. H. (2002). Acta Cryst. B58, 380–388.
- Altomare, A., Caliandro, R., Camalli, M., Cuocci, C., Giacovazzo, C., Moliterni, A. G. G. & Rizzi, R. (2004). EXPO2004. Version 2.1. http:// www.ic.cnr.it/.
- Boultif, A. & Louër, D. (2004). J. Appl. Cryst. 37, 724-731.
- Chao, M., Schempp, E. & Rosenstein, R. D. (1976). Acta Cryst. B32, 288– 290.
- Farrugia, L. J. (1997). J. Appl. Cryst. 30, 565.
- Farrugia, L. J. (1999). J. Appl. Cryst. 32, 837–838.
- Finger, L. W., Cox, D. E. & Jephcoat, A. P. (1994). J. Appl. Cryst. 27, 892– 900.
- Hanika-Heidl, H., El-din, S., Etaiw, H., Ibrahim, M. Sh., El-din, A. S. B. & Fisher, R. D. (2003). J. Organomet. Chem. 684, 329–337.
- Kuhlman, R., Schimek, G. L. & Kolis, J. W. (1999). Polyhedron, 18, 1379–1387. Larson, A. C. & Von Dreele, R. B. (2000). GSAS. Report LAUR 86-748. Los
- Alamos National Laboratory, New Mexico, USA. Le Bail, A., Duroy, H. & Fourquet, J. L. (1988). Mater. Res. Bull. 23, 447–
- 452. Macrae, C. F., Edgington, P. R., McCabe, P., Pidcock, E., Shields, G. P., Taylor,
- R., Towler, M. & van de Streek, J. (2006). J. Appl. Cryst. 39, 453–457.
- Philips (2002). X'Pert Data Collector. Version 2.0. Philips Analytical BV, Almelo, The Netherlands.
- Scott, H. G. (1983). J. Appl. Cryst. 16, 159–163.
- Stephens, P. W. (1999). J. Appl. Cryst. 32, 281–289.
- Teichert, O. & Sheldrick, W. S. (1999). Z. Anorg. Allg. Chem. 625, 1860– 1865.
- Thompson, P., Cox, D. E. & Hastings, J. B. (1987). J. Appl. Cryst. 20, 79–83.
- Tronic, T. A., deKrafft, K. E., Lim, M. J., Ley, A. N. & Pike, R. D. (2007). Inorg. Chem. 46, 8897–8912.
- Westrip, S. P. (2008). publCIF. In preparation.