

OPEN a ACCESS

# Two tautomers in the same crystal: 3-(4-fluorophenyl)-1H-pyrazole and 5-(4-fluorophenyl)-1*H*-pyrazole

#### Thammarse S. Yamuna,<sup>a</sup> Manpreet Kaur,<sup>a</sup> Jerry P. Jasinski,<sup>b\*</sup> Brian J. Anderson<sup>b</sup> and H. S. Yathirajan<sup>a</sup>

<sup>a</sup>Department of Studies in Chemistry, University of Mysore, Manasagangotri, Mysore 570 006, India, and <sup>b</sup>Department of Chemistry, Keene State College, 229 Main Street, Keene, NH 03435-2001, USA. \*Correspondence e-mail: jjasinski@keene.edu

Received 20 July 2014; accepted 23 July 2014

Edited by M. Bolte, Goethe-Universität Frankfurt, Germany

The title co-crystal, 3-(4-fluorophenyl)-1H-pyrazole-5-(4fluorophenyl)-1*H*-pyrazole (1/1), C<sub>9</sub>H<sub>7</sub>FN<sub>2</sub>, crystallizes with four independent molecules (A, B, C and D) in the asymmetric unit exhibiting two tautomeric forms (A and D; B and C) due to N-H proton exchange between the two N atoms of the pyrazole ring. The dihedral angles between the mean planes of the pyrazole and benzene rings are 15.6(1), 19.8 (9), 14.0 (1) and 10.7 (7)° in molecules A, B, C and D, respectively. In the crystal,  $N-H \cdots N$  hydrogen bonds link the four molecules in the asymmetric unit into a ring with an  $R_4^4(12)$  motif. Furthermore, weak C-H···F interactions link the molecules into a three-dimensional network.

Keywords: crystal structure; pyrazole derivative; tautomeric forms; hydrogen bonds.

#### CCDC reference: 1015543

#### 1. Related literature

For biological and pharmacological properties of pyrazole compounds, see: Isloor et al. (2009); Patel et al. (2010); Sarojini et al. (2010); Samshuddin et al. (2012). For related structures, see: Baktır et al. (2011); Fun et al. (2012); Yamuna et al. (2013). For bond-length data, see: Allen et al. (1987). For a description of hydrogen bonds, see: Etter et al. (1990).



# 2. Experimental

#### 2.1. Crystal data

$C_9H_7FN_2$	$\gamma = 68.249 \ (5)^{\circ}$
$M_r = 162.17$	$V = 1643.57 (16) \text{ Å}^3$
Triclinic, P1	Z = 8
a = 10.3961 (5)  Å	Cu $K\alpha$ radiation
b = 10.8565 (6) Å	$\mu = 0.81 \text{ mm}^{-1}$
c = 16.1431 (7) Å	$T = 173  { m K}$
$\alpha = 84.704 \ (4)^{\circ}$	$0.22 \times 0.16 \times 0.10 \text{ mm}$
$\beta = 76.223 \ (4)^{\circ}$	

#### 2.2. Data collection

Agilent Eos Gemini diffractometer Absorption correction: multi-scan (CrysAlis PRO and CrysAlis RED, Agilent (2012).  $T_{\min} = 0.881, T_{\max} = 1.000$ 

2.3. Refinement  $R[F^2 > 2\sigma(F^2)] = 0.044$ 

$wR(F^2) = 0.122$	
S = 1.03	
6209 reflections	
449 parameters	

 $R_{\rm int} = 0.019$ 

11343 measured reflections

6209 independent reflections

5042 reflections with  $I > 2\sigma(I)$ 

Table 1

Hydrogen-bond geometry (Å, °).

$D-H\cdots A$ $D-H$ $H\cdots A$ $D\cdots A$ $D-H\cdots$	
	$H \cdot \cdot \cdot A$
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	(2) (2) (2) (3)

Symmetry codes: (i) x, y - 1, z; (ii) -x + 1, -y + 2, -z + 1; (iii) x + 1, y, z.

Data collection: CrysAlis PRO (Agilent, 2012); cell refinement: CrysAlis PRO; data reduction: CrysAlis RED (Agilent, 2012); program(s) used to solve structure: SUPERFLIP (Palatinus & Chapuis, 2007); program(s) used to refine structure: SHELXL2012 (Sheldrick, 2008); molecular graphics: OLEX2 (Dolomanov et al., 2009); software used to prepare material for publication: OLEX2.

#### Acknowledgements

TSY thanks the University of Mysore for research facilities and also is grateful to the Principal, Maharani's Science College for Women, Mysore, for giving permission to undertake research. JPJ acknowledges the NSF-MRI program (grant No. CHE-1039027) for funds to purchase the X-ray diffractometer.

Supporting information for this paper is available from the IUCr electronic archives (Reference: BT6989).

Acta Cryst. E68, o2680.

#### References

- Agilent (2012). CrysAlis PRO and CrysAlis RED. Agilent Technologies, Yarnton, England.
- Allen, F. H., Kennard, O., Watson, D. G., Brammer, L., Orpen, A. G. & Taylor, R. (1987). J. Chem. Soc. Perkin Trans. 2, pp. S1–19.
- Baktır, Z., Akkurt, M., Samshuddin, S., Narayana, B. & Yathirajan, H. S. (2011). Acta Cryst. E67, o1292-o1293.
- Dolomanov, O. V., Bourhis, L. J., Gildea, R. J., Howard, J. A. K. & Puschmann, H. (2009). J. Appl. Cryst. 42, 339–341.
- Etter, M. C., MacDonald, J. C. & Bernstein, J. (1990). Acta Cryst. B46, 256–262. Fun, H.-K., Chia, T. S., Sapnakumari, M., Narayana, B. & Sarojini, B. K. (2012).
- Isloor, A. M., Kalluraya, B. & Shetty, P. (2009). Eur. J. Med. Chem. 44, 3784–3787.
- Palatinus, L. & Chapuis, G. (2007). J. Appl. Cryst. 40, 786-790.
- Patel, C. K., Rami, C. S., Panigrahi, B. & Patel, C. N. (2010). J. Chem. Pharm. Res. 2, 73–78.
- Samshuddin, S., Narayana, B., Yathirajan, H. S., Gerber, T., Hosten, E. & Betz, R. (2012). Acta Cryst. E68, 03216–03217.
- Sarojini, B. K., Vidyagayatri, M., Darshanraj, C. G., Bharath, B. R. & Manjunatha, H. (2010). Lett. Drug Des. Discov. 7, 214–224.
- Sheldrick, G. M. (2008). Acta Cryst. A64, 112-122.
- Yamuna, T. S., Jasinski, J. P., Scadova, D. R., Yathirajan, H. S. & Kaur, M. (2013). Acta Cryst. E69, 01425-01426.

# supporting information

Acta Cryst. (2014). E70, o949–o950 [doi:10.1107/S160053681401695X]

# Two tautomers in the same crystal: 3-(4-fluorophenyl)-1*H*-pyrazole and 5-(4-fluorophenyl)-1*H*-pyrazole

# Thammarse S. Yamuna, Manpreet Kaur, Jerry P. Jasinski, Brian J. Anderson and H. S. Yathirajan

# S1. Structural commentary

Pyrazoles are an important class of heterocyclic compounds and many pyrazole derivatives are reported to have a broad spectrum of biological properties, including antibacterial and anti-inflammatory activities (Patel *et al.*, 2010), anticancer (Sarojini *et al.*, 2010; Samshuddin *et al.*, 2012) anti-inflammatory, antidepressant, anticonvulsant and anti-HIV properties (Isloor *et al.*, 2009). Because of these various interesting fields of application as well as their fairly assessable path of synthesis, the pyrazoline ring became a center of attraction for organic chemists. Crystal structures of some related compounds include 3, 5-bis(4-fluorophenyl)-1-(4-nitrophenyl)-4,5-dihydro-1H-pyrazole (Samshuddin *et al.*, 2012), 5-(4-bromophenyl)- 3-(4-fluorophenyl)-1-phenyl-4,5-dihydro-1H-pyrazole (Fun *et al.*, 2012), 3,5-bis(4-fluorophenyl)-4,5-di-hydro-1H-pyrazole (Fun *et al.*, 2012), 3,5-bis(4-fluorophenyl)-4,5-di-hydro-1H-pyrazole. In trifluoroacetate (Yamuna *et al.*, 2013). In view of the importance of the title compound, C<sub>9</sub>H<sub>7</sub>FN<sub>2</sub>, the paper reports its crystal structure.

The title compound, C<sub>9</sub>H<sub>7</sub>FN<sub>2</sub>, crystallises with four independent molecules (A,B,C and D) in the asymmetric unit exhibiting two tautomeric forms (A and D; B and C) due to N—H proton exchange between the two nitrogen atoms (N1 and N2) of the pyrazole ring (Fig. 1). The dihedral angles between the mean planes of the pyrazole ring and phenyl ring are 15.6 (1)°, 19.8 (9)°, 14.0 (1)° and 10.7 (7)°, in the molecules A, B, C and D, respectively. Bond lengths are in normal ranges (Allen *et al.*, 1987). In the crystal, N—H···N intermolecular hydrogen bonds link the four molecules in the asymmetric unit to a ring with motif  $R_4^4(12)$  (Etter *et al.*, 1990). Furthermore, weak C—H···F intermolecular interactions link the molecules to a three-dimensional network (Fig. 2).

# S2. Synthesis and crystallization

Commercially available 3-(4-fluorophenyl)-1H-pyrazole was dissolved in 5 ml of dimethylformamide at 303 K over a heating magnetic stirrer. X-ray quality crystals were formed on slow evaporation. (m.p.: 368-373 K).

### **S3. Refinement**

The H atoms bonded to N (H2A, H1B, H1C and H2D) were refined isotropically and all of the remaining H atoms were placed in their calculated positions and then refined using the riding model with C—H lengths of 0.93Å. Isotropic displacement parameters for these atoms were set to 1.2 times  $U_{eq}$  of the parent C atom.



# Figure 1

ORTEP drawing of the title compound showing the labeling scheme of the asymmetric unit of the title compound with 30% probability displacement ellipsoids.



### Figure 2

Molecular packing for the title compound viewed along the *c* axis. Dashed lines indicate N—H…N intermolecular hydrogen bonds and weak C—H…F intermolecular interactions together forming a 2D supramolecular network structure. H atoms not involved in hydrogen bonding have been removed for clarity.



## Figure 3

The two tautomers in the same crystal: 3-(4-fluorophenyl)-1*H*-pyrazole (left) and 5-(4-fluorophenyl)-1*H*-pyrazole (right).

#### 3-(4-Fluorophenyl)-1*H*-pyrazole–5-(4-fluorophenyl)-1*H*-pyrazole (1/1)

Crystal data	
C <sub>9</sub> H <sub>7</sub> FN <sub>2</sub> $M_r = 162.17$ Triclinic, $P\overline{1}$ a = 10.3961 (5) Å b = 10.8565 (6) Å c = 16.1431 (7) Å $a = 84.704 (4)^{\circ}$ $\beta = 76.223 (4)^{\circ}$ $\gamma = 68.249 (5)^{\circ}$ $V = 1643.57 (16) Å^{3}$	Z = 8 F(000) = 672 $D_x = 1.311 \text{ Mg m}^{-3}$ Cu Ka radiation, $\lambda = 1.54184 \text{ Å}$ Cell parameters from 4341 reflections $\theta = 4.4-71.4^{\circ}$ $\mu = 0.81 \text{ mm}^{-1}$ T = 173  K Irregular, colourless $0.22 \times 0.16 \times 0.10 \text{ mm}$
Data collection	
Agilent Eos Gemini diffractometer Radiation source: Enhance (Cu) X-ray Source Graphite monochromator Detector resolution: 16.0416 pixels mm <sup>-1</sup> ω scans Absorption correction: multi-scan ( <i>CrysAlis PRO</i> and <i>CrysAlis RED</i> , Agilent (2012).	$T_{\min} = 0.881, T_{\max} = 1.000$ 11343 measured reflections 6209 independent reflections 5042 reflections with $I > 2\sigma(I)$ $R_{\text{int}} = 0.019$ $\theta_{\max} = 71.3^{\circ}, \theta_{\min} = 4.4^{\circ}$ $h = -12 \rightarrow 6$ $k = -13 \rightarrow 12$ $l = -19 \rightarrow 18$
Refinement	
Refinement on $F^2$ Least-squares matrix: full $R[F^2 > 2\sigma(F^2)] = 0.044$ $wR(F^2) = 0.122$ S = 1.03 6209 reflections 449 parameters 0 restraints	Primary atom site location: structure-invariant direct methods Hydrogen site location: mixed H atoms treated by a mixture of independent and constrained refinement $w = 1/[\sigma^2(F_o^2) + (0.0568P)^2 + 0.3619P]$ where $P = (F_o^2 + 2F_c^2)/3$ $(\Delta/\sigma)_{max} < 0.001$ $\Delta\rho_{max} = 0.26$ e Å <sup>-3</sup> $\Delta\rho_{min} = -0.22$ e Å <sup>-3</sup>

#### Special details

**Geometry**. All esds (except the esd in the dihedral angle between two l.s. planes) are estimated using the full covariance matrix. The cell esds are taken into account individually in the estimation of esds in distances, angles and torsion angles; correlations between esds in cell parameters are only used when they are defined by crystal symmetry. An approximate (isotropic) treatment of cell esds is used for estimating esds involving l.s. planes.

	x	У	Ζ	$U_{ m iso}$ */ $U_{ m eq}$	
F1A	0.91461 (15)	0.91870 (11)	0.36091 (9)	0.0741 (4)	
N1A	0.86295 (14)	0.36783 (13)	0.29541 (9)	0.0411 (3)	
N2A	0.89322 (16)	0.23574 (14)	0.30565 (10)	0.0464 (3)	
H2A	0.850 (3)	0.194 (3)	0.2780 (16)	0.085 (8)*	
C1A	0.93710 (16)	0.40020 (15)	0.34199 (9)	0.0368 (3)	
C2A	1.01500 (19)	0.28686 (17)	0.38175 (11)	0.0452 (4)	
H2AA	1.0764	0.2810	0.4183	0.054*	
C3A	0.9839 (2)	0.18559 (18)	0.35684 (12)	0.0493 (4)	
H3A	1.0208	0.0952	0.3733	0.059*	
C4A	0.92972 (16)	0.53758 (15)	0.34570 (9)	0.0357 (3)	
C5A	0.81711 (18)	0.64507 (17)	0.32574 (11)	0.0436 (4)	
H5A	0.7430	0.6299	0.3085	0.052*	
C6A	0.8112 (2)	0.77360 (17)	0.33054 (12)	0.0512 (4)	
H6A	0.7345	0.8467	0.3164	0.061*	
C7A	0.9187 (2)	0.79295 (17)	0.35624 (12)	0.0499 (4)	
C8A	1.0311 (2)	0.69078 (19)	0.37737 (12)	0.0521 (4)	
H8A	1.1037	0.7075	0.3954	0.063*	
C9A	1.03655 (19)	0.56220 (17)	0.37178 (11)	0.0455 (4)	
H9A	1.1140	0.4899	0.3859	0.055*	
F1B	0.3343 (2)	1.06776 (13)	0.43105 (13)	0.1157 (6)	
N1B	0.45578 (15)	0.46841 (13)	0.33291 (8)	0.0390 (3)	
H1B	0.500 (3)	0.495 (2)	0.2756 (17)	0.088 (8)*	
N2B	0.45026 (16)	0.34515 (13)	0.34547 (9)	0.0451 (3)	
C1B	0.39261 (16)	0.54133 (15)	0.40414 (9)	0.0336 (3)	
C2B	0.34397 (18)	0.46175 (16)	0.46561 (10)	0.0425 (4)	
H2B	0.2948	0.4846	0.5229	0.051*	
C3B	0.3821 (2)	0.34208 (16)	0.42597 (11)	0.0473 (4)	
H3B	0.3623	0.2674	0.4527	0.057*	
C4B	0.37974 (16)	0.67949 (15)	0.40913 (10)	0.0357 (3)	
C5B	0.3388 (2)	0.73875 (18)	0.48847 (12)	0.0514 (4)	
H5B	0.3209	0.6886	0.5384	0.062*	
C6B	0.3238 (2)	0.8687 (2)	0.49627 (16)	0.0677 (6)	
H6B	0.2963	0.9084	0.5509	0.081*	
C7B	0.3493 (3)	0.93936 (19)	0.42403 (18)	0.0711 (6)	
C8B	0.3899 (3)	0.8856 (2)	0.34485 (17)	0.0778 (7)	
H8B	0.4076	0.9369	0.2955	0.093*	
C9B	0.4048 (2)	0.75479 (19)	0.33745 (13)	0.0581 (5)	
H9B	0.4327	0.7162	0.2825	0.070*	
F1C	0.06409 (14)	0.15006 (18)	0.11668 (9)	0.0902 (5)	
N1C	0.62673 (16)	0.14637 (14)	0.21875 (9)	0.0428 (3)	
H1C	0.560 (3)	0.214 (3)	0.2606 (19)	0.106 (9)*	
N2C	0.76704 (16)	0.10753 (14)	0.21687 (10)	0.0496 (4)	
C1C	0.60268 (18)	0.07149 (15)	0.16611 (10)	0.0397 (3)	
C2C	0.7333 (2)	-0.01966 (17)	0.12914 (12)	0.0485 (4)	
H2C	0.7522	-0.0865	0.0891	0.058*	

Fractional atomic coordinates and isotropic or equivalent isotropic displacement parameters  $(\hat{A}^2)$ 

C3C	0.8310 (2)	0.00694 (18)	0.16274 (13)	0.0521 (4)
H3C	0.9306	-0.0405	0.1489	0.063*
C4C	0.46026 (18)	0.09391 (16)	0.15406 (10)	0.0402 (4)
C5C	0.4371 (2)	-0.0033 (2)	0.11510 (12)	0.0559 (5)
H5C	0.5139	-0.0835	0.0969	0.067*
C6C	0.3038 (3)	0.0154 (2)	0.10251 (14)	0.0673 (6)
H6C	0.2885	-0.0511	0.0761	0.081*
C7C	0.1955 (2)	0.1309 (3)	0.12873 (12)	0.0602 (5)
C8C	0.2129 (2)	0.2291 (2)	0.16722 (12)	0.0581 (5)
H8C	0.1350	0.3088	0.1849	0.070*
C9C	0.34629 (18)	0.21022 (18)	0.17996 (11)	0.0470 (4)
H9C	0.3597	0.2776	0.2067	0.056*
F1D	0.00377 (12)	0.72341 (13)	0.04635 (8)	0.0704 (3)
N1D	0.59139 (15)	0.52634 (14)	0.16671 (8)	0.0429 (3)
N2D	0.72929 (16)	0.51037 (17)	0.15930 (10)	0.0501 (4)
H2D	0.778 (3)	0.459 (3)	0.206 (2)	0.114 (10)*
C1D	0.55077 (17)	0.59754 (16)	0.09872 (9)	0.0388 (3)
C2D	0.6648 (2)	0.62687 (19)	0.04769 (11)	0.0503 (4)
H2DA	0.6660	0.6760	-0.0042	0.060*
C3D	0.7748 (2)	0.5694 (2)	0.08865 (12)	0.0562 (5)
H3D	0.8677	0.5715	0.0697	0.067*
C4D	0.40598 (17)	0.63199 (15)	0.08592 (10)	0.0380 (3)
C5D	0.29879 (19)	0.61209 (19)	0.14987 (11)	0.0507 (4)
H5D	0.3192	0.5763	0.2032	0.061*
C6D	0.1633 (2)	0.6436 (2)	0.13699 (12)	0.0570 (5)
H6D	0.0906	0.6300	0.1808	0.068*
C7D	0.13651 (18)	0.69495 (18)	0.05921 (12)	0.0489 (4)
C8D	0.2375 (2)	0.71705 (17)	-0.00511 (11)	0.0492 (4)
H8D	0.2156	0.7536	-0.0580	0.059*
C9D	0.37280 (19)	0.68496 (16)	0.00855 (10)	0.0436 (4)
H9D	0.4443	0.6994	-0.0358	0.052*

Atomic displacement parameters  $(Å^2)$ 

	$U^{11}$	$U^{22}$	$U^{33}$	$U^{12}$	$U^{13}$	$U^{23}$
F1A	0.0908 (9)	0.0394 (6)	0.0995 (10)	-0.0298 (6)	-0.0218 (7)	-0.0071 (6)
N1A	0.0438 (7)	0.0414 (7)	0.0421 (7)	-0.0192 (6)	-0.0082 (6)	-0.0062 (6)
N2A	0.0523 (8)	0.0424 (8)	0.0506 (8)	-0.0241 (7)	-0.0091 (7)	-0.0069 (6)
C1A	0.0385 (8)	0.0405 (8)	0.0324 (7)	-0.0167 (6)	-0.0045 (6)	-0.0034 (6)
C2A	0.0554 (10)	0.0409 (9)	0.0451 (9)	-0.0210 (8)	-0.0162 (8)	0.0013 (7)
C3A	0.0580 (10)	0.0389 (9)	0.0532 (10)	-0.0202 (8)	-0.0123 (8)	0.0006 (7)
C4A	0.0391 (8)	0.0377 (8)	0.0299 (7)	-0.0159 (6)	-0.0023 (6)	-0.0026 (6)
C5A	0.0448 (9)	0.0439 (9)	0.0440 (9)	-0.0180 (7)	-0.0096 (7)	-0.0004 (7)
C6A	0.0542 (10)	0.0386 (9)	0.0566 (11)	-0.0124 (8)	-0.0119 (8)	0.0013 (8)
C7A	0.0627 (11)	0.0362 (9)	0.0532 (10)	-0.0229 (8)	-0.0068 (8)	-0.0061 (7)
C8A	0.0553 (10)	0.0502 (10)	0.0607 (11)	-0.0263 (8)	-0.0163 (9)	-0.0072 (8)
C9A	0.0469 (9)	0.0413 (9)	0.0521 (10)	-0.0164 (7)	-0.0159 (8)	-0.0032 (7)
F1B	0.1591 (16)	0.0346 (7)	0.1658 (17)	-0.0410 (9)	-0.0453 (13)	-0.0091 (8)

N1B	0.0529 (8)	0.0327 (6)	0.0319 (7)	-0.0170 (6)	-0.0067 (6)	-0.0028 (5)
N2B	0.0587 (9)	0.0327 (7)	0.0467 (8)	-0.0178 (6)	-0.0127 (7)	-0.0054 (6)
C1B	0.0371 (8)	0.0331 (7)	0.0321 (7)	-0.0139 (6)	-0.0083 (6)	0.0000 (6)
C2B	0.0522 (10)	0.0391 (8)	0.0366 (8)	-0.0214 (7)	-0.0028 (7)	0.0004 (6)
C3B	0.0612 (11)	0.0357 (8)	0.0517 (10)	-0.0265 (8)	-0.0122 (8)	0.0046 (7)
C4B	0.0358 (8)	0.0323 (7)	0.0391 (8)	-0.0128 (6)	-0.0071 (6)	-0.0011 (6)
C5B	0.0625 (11)	0.0395 (9)	0.0488 (10)	-0.0170 (8)	-0.0049 (8)	-0.0092 (7)
C6B	0.0804 (15)	0.0426 (10)	0.0767 (14)	-0.0170 (10)	-0.0115 (11)	-0.0216 (10)
C7B	0.0859 (15)	0.0292 (9)	0.1048 (18)	-0.0217 (9)	-0.0287 (14)	-0.0076 (10)
C8B	0.119 (2)	0.0464 (11)	0.0788 (16)	-0.0452 (13)	-0.0219 (14)	0.0153 (11)
C9B	0.0885 (15)	0.0437 (10)	0.0467 (10)	-0.0342 (10)	-0.0078 (10)	0.0029 (8)
F1C	0.0624 (8)	0.1541 (15)	0.0731 (9)	-0.0566 (9)	-0.0189 (7)	-0.0068 (9)
N1C	0.0485 (8)	0.0366 (7)	0.0464 (8)	-0.0168 (6)	-0.0111 (6)	-0.0065 (6)
N2C	0.0511 (8)	0.0389 (7)	0.0637 (9)	-0.0168 (6)	-0.0203 (7)	-0.0021 (7)
C1C	0.0533 (9)	0.0337 (8)	0.0354 (8)	-0.0190 (7)	-0.0100 (7)	-0.0012 (6)
C2C	0.0563 (10)	0.0386 (9)	0.0508 (10)	-0.0135 (8)	-0.0154 (8)	-0.0082 (7)
C3C	0.0493 (10)	0.0410 (9)	0.0638 (11)	-0.0101 (8)	-0.0171 (9)	-0.0031 (8)
C4C	0.0501 (9)	0.0444 (9)	0.0302 (7)	-0.0236 (7)	-0.0059 (7)	0.0001 (6)
C5C	0.0648 (12)	0.0559 (11)	0.0554 (11)	-0.0272 (9)	-0.0157 (9)	-0.0112 (9)
C6C	0.0776 (15)	0.0851 (16)	0.0592 (12)	-0.0482 (13)	-0.0175 (11)	-0.0111 (11)
C7C	0.0541 (11)	0.0995 (17)	0.0409 (9)	-0.0436 (11)	-0.0104 (8)	0.0011 (10)
C8C	0.0469 (10)	0.0782 (14)	0.0454 (10)	-0.0226 (9)	-0.0005 (8)	-0.0070 (9)
C9C	0.0480 (9)	0.0548 (10)	0.0393 (9)	-0.0221 (8)	-0.0035 (7)	-0.0069 (7)
F1D	0.0458 (6)	0.0818 (9)	0.0781 (8)	-0.0133 (6)	-0.0200 (6)	0.0014 (7)
N1D	0.0442 (7)	0.0503 (8)	0.0321 (7)	-0.0153 (6)	-0.0061 (6)	-0.0040 (6)
N2D	0.0475 (8)	0.0618 (10)	0.0423 (8)	-0.0202 (7)	-0.0096 (7)	-0.0047 (7)
C1D	0.0461 (9)	0.0385 (8)	0.0300 (7)	-0.0159 (7)	-0.0027 (6)	-0.0039 (6)
C2D	0.0540 (10)	0.0567 (11)	0.0419 (9)	-0.0266 (9)	-0.0046 (8)	0.0042 (8)
C3D	0.0503 (10)	0.0704 (13)	0.0528 (11)	-0.0311 (9)	-0.0039 (8)	-0.0038 (9)
C4D	0.0449 (8)	0.0335 (7)	0.0323 (7)	-0.0130 (6)	-0.0036 (6)	-0.0025 (6)
C5D	0.0481 (10)	0.0642 (11)	0.0343 (8)	-0.0183 (8)	-0.0049 (7)	0.0061 (8)
C6D	0.0452 (10)	0.0727 (13)	0.0454 (10)	-0.0199 (9)	0.0016 (8)	0.0012 (9)
C7D	0.0409 (9)	0.0461 (9)	0.0546 (10)	-0.0078 (7)	-0.0125 (8)	-0.0047 (8)
C8D	0.0583 (11)	0.0444 (9)	0.0420 (9)	-0.0140 (8)	-0.0151 (8)	0.0044 (7)
C9D	0.0512 (9)	0.0420 (9)	0.0359 (8)	-0.0175 (7)	-0.0059 (7)	0.0023 (7)

Geometric parameters (Å, °)

F1A—C7A	1.3584 (19)	F1C—C7C	1.362 (2)
N1A—N2A	1.3525 (19)	N1C—H1C	0.98 (3)
N1A—C1A	1.345 (2)	N1C—N2C	1.353 (2)
N2A—H2A	0.94 (3)	N1C—C1C	1.351 (2)
N2A—C3A	1.333 (2)	N2C—C3C	1.326 (2)
C1A—C2A	1.395 (2)	C1C—C2C	1.381 (2)
C1A—C4A	1.471 (2)	C1C—C4C	1.466 (2)
C2A—H2AA	0.9500	C2C—H2C	0.9500
C2A—C3A	1.372 (2)	C2C—C3C	1.384 (3)
СЗА—НЗА	0.9500	СЗС—НЗС	0.9500

C4A—C5A	1.391 (2)	C4C—C5C	1.396 (2)
С4А—С9А	1.395 (2)	C4C—C9C	1.388 (2)
С5А—Н5А	0.9500	C5C—H5C	0.9500
C5A—C6A	1.382 (2)	C5C—C6C	1.386 (3)
С6А—Н6А	0.9500	C6C—H6C	0.9500
C6A—C7A	1.370 (3)	C6C—C7C	1.357 (3)
C7A—C8A	1.368 (3)	C7C—C8C	1.369 (3)
С8А—Н8А	0.9500	C8C—H8C	0.9500
C8A—C9A	1 387 (2)	C8C - C9C	1 387 (3)
С9А—Н9А	0.9500	C9C—H9C	0.9500
F1B—C7B	1 356 (2)	F1D-C7D	1 361 (2)
N1B—H1B	1.00(2)	N1D-N2D	1.301(2) 1.356(2)
N1B—N2B	1.00(5) 1.3551(18)	NID-CID	1.333(2)
N1B-C1B	1.3331(10) 1.343(2)	N2D—H2D	1.07(2)
N2B_C3B	1.343(2) 1.328(2)	N2D_C3D	1.02(5)
C1B-C2B	1.320(2) 1.383(2)	C1D-C2D	1.327(2) 1.397(2)
C1B - C2B	1.365(2)	C1D - C2D	1.397(2) 1.471(2)
$C_{1D}$ $C_{4D}$ $C_{2D}$ $U_{2D}$	0.0500	C1D - C4D	1.471 (2)
$C_{2D}$ $C_{2D}$ $C_{2D}$	0.9300	C2D— $H2DAC2D$ — $C2D$	0.9300
C2D U2D	1.360 (2)	C2D = C3D	1.575 (5)
	0.9300	C3D—H3D C4D—C5D	0.9300
C4B = C3B	1.308(2) 1.384(2)	C4D = C3D	1.394(2)
C4B—C9B	1.384 (2)	C4D—C9D	1.391 (2)
C3B—H3B	0.9500	CSD—HSD	0.9500
CSB-C6B	1.375 (3)	CSD—C6D	1.383 (3)
С6В—Н6В	0.9500	C6D—H6D	0.9500
C6B—C/B	1.362 (3)	C6D—C/D	1.374 (3)
C/B—C8B	1.361 (3)	C/D—C8D	1.364 (3)
C8B—H8B	0.9500	C8D—H8D	0.9500
C8B—C9B	1.383 (3)	C8D—C9D	1.385 (2)
С9В—Н9В	0.9500	C9D—H9D	0.9500
C1A—N1A—N2A	106.39 (13)	N2C—N1C—H1C	118.1 (16)
N1A—N2A—H2A	118.9 (16)	C1C—N1C—H1C	130.5 (17)
C3A—N2A—N1A	110.49 (14)	C1C—N1C—N2C	110.98 (14)
C3A—N2A—H2A	130.6 (16)	C3C—N2C—N1C	105.89 (14)
N1A—C1A—C2A	109.52 (14)	N1C—C1C—C2C	106.74 (15)
N1A—C1A—C4A	121.34 (14)	N1C—C1C—C4C	122.62 (15)
C2A—C1A—C4A	129.14 (14)	C2C—C1C—C4C	130.62 (15)
C1A—C2A—H2AA	127.4	C1C—C2C—H2C	127.3
C3A—C2A—C1A	105.28 (15)	C1C—C2C—C3C	105.42 (15)
СЗА—С2А—Н2АА	127.4	C3C—C2C—H2C	127.3
N2A—C3A—C2A	108.33 (16)	N2C—C3C—C2C	110.97 (16)
N2A—C3A—H3A	125.8	N2C—C3C—H3C	124.5
С2А—С3А—Н3А	125.8	С2С—С3С—Н3С	124.5
C5A—C4A—C1A	121.94 (14)	C5C—C4C—C1C	119.78 (16)
C5A—C4A—C9A	118.47 (15)	C9C—C4C—C1C	121.88 (15)
C9A—C4A—C1A	119.57 (14)	C9C—C4C—C5C	118.33 (17)
С4А—С5А—Н5А	119.4	C4C—C5C—H5C	119.5

C6A—C5A—C4A	121.12 (16)	C6C—C5C—C4C	121.1 (2)
С6А—С5А—Н5А	119.4	C6C—C5C—H5C	119.5
С5А—С6А—Н6А	120.8	С5С—С6С—Н6С	120.7
C7A—C6A—C5A	118.34 (17)	C7C—C6C—C5C	118.55 (19)
С7А—С6А—Н6А	120.8	С7С—С6С—Н6С	120.7
F1A—C7A—C6A	118.98 (17)	F1C—C7C—C8C	118.5 (2)
F1A—C7A—C8A	118.17 (17)	C6C—C7C—F1C	118.9 (2)
C8A—C7A—C6A	122.85 (16)	C6C—C7C—C8C	122.57 (18)
С7А—С8А—Н8А	120.8	C7C—C8C—H8C	120.6
C7A—C8A—C9A	118.35 (17)	C7C—C8C—C9C	118.84 (19)
С9А—С8А—Н8А	120.8	C9C—C8C—H8C	120.6
C4A—C9A—H9A	119.6	C4C-C9C-H9C	119.7
C8A - C9A - C4A	120.86 (16)	C8C - C9C - C4C	120.64(17)
C8A - C9A - H9A	119.6	C8C - C9C - H9C	1197
N2B-N1B-H1B	120.0 (15)	C1D - N1D - N2D	106 73 (14)
C1B—N1B—H1B	128.9 (15)	N1D - N2D - H2D	117.9(17)
C1B $N1B$ $N2B$	111.08(13)	C3D N2D N1D	117.5(17)
C3B = N2B = N1B	105 72 (13)	C3D = N2D = H2D	1319(17)
N1B-C1B-C2B	106.85(14)	N1D - C1D - C2D	109.36(17)
NIB CIB C/B	100.03(14) 123 31(13)	N1D = C1D = C4D	109.30(13) 121.20(14)
$C^{2}$ $C^{1}$ $C^{4}$ $C^{4$	123.31(13) 129.83(14)	$C^{2}D$ $C^{1}D$ $C^{4}D$	121.20(14) 120.44(15)
$C_{2D}$ $C_{1D}$ $C_{4D}$ $C_{4D}$	129.85 (14)	C1D $C2D$ $H2DA$	129.44 (13)
$C_{1D}$ $-C_{2D}$ $-C_{12D}$	127.3 105.27 (14)	C1D = C2D = C1D	127.4
$C_{2}D = C_{2}D = U_{2}D$	103.37 (14)	$C_{2D}$ $C_{2D}$ $C_{2D}$ $C_{2D}$ $C_{2D}$	103.10 (10)
$C_{3}D - C_{2}D - \Pi_{2}D$	127.5	$C_{2D}$ $C_{2D}$ $C_{2D}$ $C_{2D}$	127.4
N2B-C3B-C2B	110.99 (14)	N2D = C3D = U2D	108.57 (16)
N2B - C3B - H3B	124.5	$N_2D = C_3D = H_3D$	125.7
$C_{2B}$ — $C_{3B}$ — $H_{3B}$	124.5	C2D = C3D = H3D	125.7
$C_{3B}$ $C_{4B}$ $C_{1B}$	119.17 (14)	CSD—C4D—C1D	121.57 (15)
	122.57 (15)	C9D—C4D—C1D	120.35 (14)
C9B—C4B—C5B	118.26 (16)	C9D—C4D—C5D	118.08 (16)
C4B—C5B—H5B	119.4	C4D—C5D—H5D	119.4
C6B—C5B—C4B	121.24 (18)	C6D—C5D—C4D	121.17 (16)
С6В—С5В—Н5В	119.4	C6D—C5D—H5D	119.4
С5В—С6В—Н6В	120.7	C5D—C6D—H6D	120.8
C7B—C6B—C5B	118.6 (2)	C7D—C6D—C5D	118.35 (17)
С7В—С6В—Н6В	120.7	C7D—C6D—H6D	120.8
F1B—C7B—C6B	119.0 (2)	F1D—C7D—C6D	118.09 (17)
F1B—C7B—C8B	118.7 (2)	F1D—C7D—C8D	119.25 (16)
C8B—C7B—C6B	122.33 (18)	C8D—C7D—C6D	122.65 (17)
C7B—C8B—H8B	120.6	C7D—C8D—H8D	120.8
C7B—C8B—C9B	118.8 (2)	C7D—C8D—C9D	118.44 (16)
C9B—C8B—H8B	120.6	C9D—C8D—H8D	120.8
C4B—C9B—H9B	119.6	C4D—C9D—H9D	119.3
C8B—C9B—C4B	120.79 (19)	C8D—C9D—C4D	121.31 (16)
C8B—C9B—H9B	119.6	C8D—C9D—H9D	119.3
F1A—C7A—C8A—C9A	-179.30 (16)	F1C—C7C—C8C—C9C	179.81 (17)
N1A—N2A—C3A—C2A	0.2 (2)	N1C—N2C—C3C—C2C	0.2 (2)

N1A—C1A—C2A—C3A	-0.03 (19)	N1C—C1C—C2C—C3C	-0.09 (19)
N1A—C1A—C4A—C5A	20.9 (2)	N1C—C1C—C4C—C5C	-165.31 (16)
N1A—C1A—C4A—C9A	-160.61 (15)	N1C-C1C-C4C-C9C	15.0 (2)
N2A—N1A—C1A—C2A	0.12 (18)	N2C—N1C—C1C—C2C	0.23 (19)
N2A—N1A—C1A—C4A	179.93 (13)	N2C—N1C—C1C—C4C	-178.51 (14)
C1A—N1A—N2A—C3A	-0.17 (19)	C1C—N1C—N2C—C3C	-0.28 (19)
C1A—C2A—C3A—N2A	-0.1 (2)	C1C—C2C—C3C—N2C	-0.1 (2)
C1A—C4A—C5A—C6A	179.25 (15)	C1C—C4C—C5C—C6C	-179.71 (17)
C1A—C4A—C9A—C8A	-178.84 (16)	C1C—C4C—C9C—C8C	179.57 (16)
C2A—C1A—C4A—C5A	-159.34 (17)	C2C—C1C—C4C—C5C	16.3 (3)
C2A—C1A—C4A—C9A	19.2 (2)	C2C—C1C—C4C—C9C	-163.36 (18)
C4A—C1A—C2A—C3A	-179.82 (16)	C4C—C1C—C2C—C3C	178.51 (17)
C4A—C5A—C6A—C7A	-0.5 (3)	C4C—C5C—C6C—C7C	0.2 (3)
C5A—C4A—C9A—C8A	-0.3 (2)	C5C—C4C—C9C—C8C	-0.1 (3)
C5A—C6A—C7A—F1A	179.73 (16)	C5C—C6C—C7C—F1C	-179.94 (18)
C5A—C6A—C7A—C8A	-0.1 (3)	C5C—C6C—C7C—C8C	-0.2 (3)
C6A—C7A—C8A—C9A	0.5 (3)	C6C—C7C—C8C—C9C	0.0 (3)
C7A—C8A—C9A—C4A	-0.3 (3)	C7C—C8C—C9C—C4C	0.1 (3)
C9A—C4A—C5A—C6A	0.7 (2)	C9C—C4C—C5C—C6C	0.0 (3)
F1B-C7B-C8B-C9B	-179.8 (2)	F1D-C7D-C8D-C9D	-178.71 (16)
N1B—N2B—C3B—C2B	-0.2 (2)	N1D—N2D—C3D—C2D	0.2 (2)
N1B-C1B-C2B-C3B	-0.10 (19)	N1D—C1D—C2D—C3D	0.1 (2)
N1B—C1B—C4B—C5B	-167.30 (16)	N1D—C1D—C4D—C5D	10.9 (2)
N1B-C1B-C4B-C9B	13.7 (3)	N1D-C1D-C4D-C9D	-168.65 (15)
N2B—N1B—C1B—C2B	0.01 (18)	N2D—N1D—C1D—C2D	0.01 (19)
N2B—N1B—C1B—C4B	-178.70 (13)	N2D—N1D—C1D—C4D	179.25 (14)
C1B—N1B—N2B—C3B	0.09 (19)	C1D—N1D—N2D—C3D	-0.1 (2)
C1B—C2B—C3B—N2B	0.2 (2)	C1D—C2D—C3D—N2D	-0.2 (2)
C1B—C4B—C5B—C6B	-179.28 (18)	C1D-C4D-C5D-C6D	-179.36 (17)
C1B—C4B—C9B—C8B	179.22 (19)	C1D-C4D-C9D-C8D	179.45 (15)
C2B—C1B—C4B—C5B	14.3 (3)	C2D-C1D-C4D-C5D	-170.03 (18)
C2B—C1B—C4B—C9B	-164.73 (18)	C2D-C1D-C4D-C9D	10.4 (3)
C4B—C1B—C2B—C3B	178.49 (16)	C4D—C1D—C2D—C3D	-179.03 (16)
C4B—C5B—C6B—C7B	0.3 (3)	C4D-C5D-C6D-C7D	0.1 (3)
C5B—C4B—C9B—C8B	0.2 (3)	C5D-C4D-C9D-C8D	-0.1 (2)
C5B—C6B—C7B—F1B	179.7 (2)	C5D-C6D-C7D-F1D	178.79 (17)
C5B—C6B—C7B—C8B	-0.4 (4)	C5D-C6D-C7D-C8D	-0.6 (3)
C6B—C7B—C8B—C9B	0.4 (4)	C6D-C7D-C8D-C9D	0.7 (3)
C7B—C8B—C9B—C4B	-0.3 (4)	C7D-C8D-C9D-C4D	-0.3 (3)
C9B—C4B—C5B—C6B	-0.2 (3)	C9D—C4D—C5D—C6D	0.2 (3)

# Hydrogen-bond geometry (Å, °)

<i>D</i> —H··· <i>A</i>	D—H	H···A	D···A	<i>D</i> —H··· <i>A</i>
N2A—H2A…N2C	0.94 (3)	1.94 (3)	2.886 (2)	177 (2)
$C3A$ — $H3A$ ···F1 $A^{i}$	0.95	2.58	3.226 (2)	125
N1 <i>B</i> —H1 <i>B</i> ···N1 <i>D</i>	1.00 (3)	1.86 (3)	2.8506 (19)	175 (2)
$C3B$ — $H3B$ ···F1 $B^{i}$	0.95	2.36	3.187 (2)	145

# supporting information

$C6B$ — $H6B$ ···· $F1A^{ii}$	0.95	2.51	3.287 (3)	139
N1 <i>C</i> —H1 <i>C</i> ···N2 <i>B</i>	0.98 (3)	1.90 (3)	2.881 (2)	173 (2)
N2D— $H2D$ ··· $N1A$	1.02 (3)	1.87 (3)	2.896 (2)	178 (3)
C3D— $H3D$ ····F1 $D$ <sup>iii</sup>	0.95	2.49	3.301 (2)	143

Symmetry codes: (i) *x*, *y*–1, *z*; (ii) –*x*+1, –*y*+2, –*z*+1; (iii) *x*+1, *y*, *z*.