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3-Amino-1H-pyrazol-2-ium trifluoroacetate

T. S. Yamuna,^a Jerry P. Jasinski,^{b*} Derek R. Scadova,^b H. S. Yathirajan^a and Manpreet Kaur^a

^aDepartment of Studies in Chemistry, University of Mysore, Manasagangotri, Mysore 570 006, India, and ^bDepartment of Chemistry, Keene State College, 229 Main Street, Keene, NH 03435-2001, USA Correspondence e-mail: jjasinski@keene.edu

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Key indicators: single-crystal X-ray study; T = 173 K; mean σ (C–C) = 0.003 Å; disorder in main residue; R factor = 0.048; wR factor = 0.136; data-to-parameter ratio = 9.0.

The asymmetric unit of the title salt, $C_3H_6N_3^+ \cdot C_2F_3O_2^-$, contains two independent 3-aminopyrazolium cations and two independent trifluoroacetate anions. The F atoms of both anions were refined as disordered over two sets of sites, with common occupancy ratios of 0.639 (12):0.361 (12). In the crystal, the cations and anions are linked via N-H···O hydrogen bonds, forming chains along [100] and [010].

Related literature

For biological properties of pyrazole derivatives, see: Hall et al. (2008); Isloor et al. (2009); Patel et al. (2010); Samshuddin et al. (2010). For the chemistry of aminopyrazoles, see: Giuseppe et al. (1991). For the medicinal activity of pyrazoles, see: Vinogradov et al. (1994). For related structures, see: Dobson & Gerkin (1998); Foces-Foces et al. (1996); Hemamalini & Fun (2010); Thanigaimani et al. (2012). For hydrogen-bond graphset motifs, see: Bernstein et al. (1995). For standard bond lengths, see: Allen et al. (1987).



Experimental

Crystal data

| $C_{3}H_{6}N_{3}^{+}C_{2}F_{3}O_{2}^{-}$ |
|------------------------------------------|
| $M_r = 197.13$ |
| Monoclinic, $P2_1/n$ |
| a = 10.9292 (8) Å |
| b = 10.9332 (6) Å |
| c = 13.7002 (13) Å |
| $\beta = 107.939 \ (9)^{\circ}$ |

V = 1557.5 (2) Å³ Z = 8Cu Ka radiation $\mu = 1.58 \text{ mm}^{-1}$ T = 173 K $0.16 \times 0.14 \times 0.06 \; \rm mm$ 9227 measured reflections

 $R_{\rm int} = 0.030$

3031 independent reflections

2343 reflections with $I > 2\sigma(I)$

Data collection

Agilent Xcalibur (Eos, Gemini) diffractometer Absorption correction: multi-scan (CrysAlis PRO and CrysAlis RED; Agilent, 2012) $T_{\min} = 0.662, \ T_{\max} = 1.000$

Refinement

| $R[F^2 > 2\sigma(F^2)] = 0.048$ | 338 parameters |
|---------------------------------|------------------------------------------------------------|
| $wR(F^2) = 0.136$ | All H-atom parameters refined |
| S = 1.05 | $\Delta \rho_{\rm max} = 0.24 \text{ e } \text{\AA}^{-3}$ |
| 3031 reflections | $\Delta \rho_{\rm min} = -0.23 \ {\rm e} \ {\rm \AA}^{-3}$ |

Table 1 Hydrogen-bond geometry (Å, °).

| , , , | | | | |
|-------------------------------|----------|-------------------------|--------------|--------------------------------------|
| $D - H \cdot \cdot \cdot A$ | D-H | $H \cdot \cdot \cdot A$ | $D \cdots A$ | $D - \mathbf{H} \cdot \cdot \cdot A$ |
| $N1A - H1AA \cdots O1A^{i}$ | 0.85 (3) | 2.28 (3) | 2.936 (3) | 134 (2) |
| $N1A - H1AB \cdots O2A^{ii}$ | 0.91 (3) | 1.99 (3) | 2.884 (3) | 169 (3) |
| $N2A - H2AA \cdots O1A^{ii}$ | 0.94 (3) | 1.85 (3) | 2.778 (2) | 171 (3) |
| $N3A - H3AA \cdots O2A$ | 0.93 (3) | 1.78 (3) | 2.705 (2) | 172 (3) |
| $N1B - H1BA \cdots O2B^{iii}$ | 0.84 (3) | 2.18 (3) | 2.962 (2) | 153 (2) |
| $N1B - H1BB \cdots O2B^{iv}$ | 0.90 (3) | 2.03 (3) | 2.929 (3) | 173 (2) |
| $N2B - H2BA \cdots O1B^{iv}$ | 0.95 (3) | 1.81 (3) | 2.756 (2) | 174 (2) |
| $N3B - H3BA \cdots O1B^{v}$ | 0.91 (3) | 1.82 (3) | 2.728 (2) | 171 (2) |
| | | | | |

Symmetry codes: (i) x, y + 1, z; (ii) $-x + \frac{3}{2}, y + \frac{1}{2}, -z + \frac{1}{2}$; (iii) -x + 1, -y + 1, -z + 1; (iv) x, y - 1, z; (v) -x, -y + 1, -z + 1.

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.

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Supplementary data and figures for this paper are available from the IUCr electronic archives (Reference: LH5637).

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3-Amino-1H-pyrazol-2-ium trifluoroacetate

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S1. Comment

Pyrazoles are an important class of heterocyclic compounds and many pyrazole derivatives are reported to have a broad spectrum of biological properties, e.g. antibacterial and anti-inflammatory activities (Patel *et al.*, 2010), anticancer (Hall *et al.*, 2008), antimicrobial (Samshuddin *et al.*, 2010), anti-inflammatory, antidepressant, anticonvulsant and anti-HIV properties (Isloor *et al.*, 2009). The chemistry of aminopyrazoles has been extensively investigated in the past (Giuseppe *et al.*, 1991). The considerable biological and medicinal activities of pyrazoles (Vinogradov *et al.*, 1994) for which aminopyrazoles are preferred precursors, have stimulated our investigations.

The crystal structures of some related compounds, viz., 3-aminopyrazole-4-carboxylic acid (Dobson & Gerkin, 1998), 4-(3,5-dimethylpyrazol-1-yl)benzoic acid trifluoroacetate (Foces-Foces *et al.*, 1996), 2-amino-5-methylpyridinium trifluoroacetate (Thanigaimani *et al.*, 2012) and 2-amino-5-chloropyridinium trifluoroacetate (Hemamalini & Fun, 2010) have been reported. In view of the importance of the title compound this paper reports its crystal structure.

The asymmetric unit of the title compound consists of two crystallographically independent 3-aminopyrazolium cations (A and B) and two trifluoroacetate anions (A and B) (Fig. 1). Each 3-aminopyrazolium cation is planar, with a maximum deviation of 0.0006 (2) Å for atom N2A in cation A and 0.0005 (2) Å for atom N2B in cation B. In the cations, atoms N3A and N3B are protonated. The F atoms of both anions are disordered over two sets of positions, with occupancy ratios of 0.639 (12):0.361 (12). Bond lengths and are normal (Allen *et al.*, 1987).

In the crystal packing (Fig. 2), the A/B type 3-aminopyrazolium cations interact with the carboxylate groups of the A/B type trifluoroacetate anions through N—H···O hydrogen bonds, forming $R_2^2(8)$, $R_2^4(8)$, $R_2^4(10)$, $R_4^4(16)$ and $R_4^4(18)$ (Bernstein *et al.*, 1995) ring motifs.

S2. Experimental

A mixture of commercially available 3-aminopyrazole and trifluoroacetic acid (1:3 v/v) were stirred for 15 minutes at room temperature. X-ray quality crystals were formed on slow evaporation. (m.p.: 463-468 K).

S3. Refinement

All H atoms were located in a difference Fourier map and refined independently with isotropic displacement parameters [N-H = 0.84 (3)-0.95 (3) Å and C-H = 0.89 (3)-0.96 (3) Å].



Figure 1

The asymmetric unit of the title compound. Displacement ellipsoids are drawn at the 30% probability level. All disorder components are shown.



Figure 2

The crystal packing of the title compound, showing the hydrogen bonds (dashed lines) forming chains along [100] and [010]. H atoms not involved in hydrogen bonding and the minor component of disorder have been removed for clarity.

3-Amino-1H-pyrazol-2-ium trifluoroacetate

Crystal data

 $C_{3}H_{6}N_{3}^{+}C_{2}F_{3}O_{2}^{-}$ $M_{r} = 197.13$ Monoclinic, $P2_{1}/n$ a = 10.9292 (8) Å b = 10.9332 (6) Å c = 13.7002 (13) Å $\beta = 107.939$ (9)° V = 1557.5 (2) Å³ Z = 8

Data collection

| Bull concerton | |
|------------------------------------------------------|---------------------------------------------------------------------|
| Agilent Xcalibur (Eos, Gemini) diffractometer | $T_{\min} = 0.662, T_{\max} = 1.000$ 9227 measured reflections |
| Radiation source: Enhance (Cu) X-ray Source | 3031 independent reflections |
| Graphite monochromator | 2343 reflections with $I > 2\sigma(I)$ |
| Detector resolution: 16.0416 pixels mm ⁻¹ | $R_{\rm int} = 0.030$ |
| ω scans | $\theta_{\rm max} = 72.5^{\circ}, \ \theta_{\rm min} = 4.6^{\circ}$ |
| Absorption correction: multi-scan | $h = -13 \rightarrow 13$ |
| (CrysAlis PRO and CrysAlis RED; Agilent, | $k = -13 \rightarrow 9$ |
| 2012) | $l = -15 \rightarrow 16$ |
| Refinement | |
| Refinement on F^2 | Primary atom site location: inferred from |
| Least-squares matrix: full | neighbouring sites |
| $R[F^2 > 2\sigma(F^2)] = 0.048$ | Hydrogen site location: difference Fourier map |
| $wR(F^2) = 0.136$ | All H-atom parameters refined |
| S = 1.05 | $w = 1/[\sigma^2(F_0^2) + (0.0697P)^2 + 0.4559P]$ |
| 3031 reflections | where $P = (F_o^2 + 2F_c^2)/3$ |
| 338 parameters | $(\Delta/\sigma)_{\rm max} < 0.001$ |
| 0 restraints | $\Delta ho_{ m max} = 0.24 \ { m e} \ { m \AA}^{-3}$ |
| | $\Delta ho_{ m min} = -0.23 \ m e \ { m \AA}^{-3}$ |

F(000) = 800

 $\theta = 3.4 - 72.4^{\circ}$

 $\mu = 1.58 \text{ mm}^{-1}$

Irregular, colourless

 $0.16 \times 0.14 \times 0.06 \text{ mm}$

T = 173 K

 $D_{\rm x} = 1.681 \text{ Mg m}^{-3}$

Cu K α radiation, $\lambda = 1.5418$ Å Cell parameters from 2544 reflections

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.

Fractional atomic coordinates and isotropic or equivalent isotropic displacement parameters $(Å^2)$

| | x | у | Ζ | $U_{ m iso}$ */ $U_{ m eq}$ | Occ. (<1) |
|------|--------------|--------------|--------------|-----------------------------|-----------|
| C1A | 0.56299 (19) | 0.93721 (18) | 0.25897 (17) | 0.0384 (5) | |
| C2A | 0.4240 (2) | 0.7896 (2) | 0.25222 (19) | 0.0453 (5) | |
| H2A | 0.351 (2) | 0.741 (2) | 0.2498 (18) | 0.050 (7)* | |
| C3A | 0.4396 (2) | 0.9121 (2) | 0.26406 (18) | 0.0428 (5) | |
| H3A | 0.379 (2) | 0.968 (2) | 0.2737 (18) | 0.050 (7)* | |
| N1A | 0.6272 (2) | 1.04305 (18) | 0.2644 (2) | 0.0588 (6) | |
| H1AA | 0.593 (3) | 1.108 (3) | 0.278 (2) | 0.057 (8)* | |
| H1AB | 0.713 (3) | 1.041 (3) | 0.272 (2) | 0.064 (8)* | |
| N2A | 0.61502 (17) | 0.83064 (15) | 0.24439 (15) | 0.0396 (4) | |

| H2AA | 0.697 (3) | 0.815 (3) | 0.240 (2) | 0.062 (8)* | |
|------|--------------|--------------|--------------|-------------|------------|
| N3A | 0.52875 (17) | 0.73971 (17) | 0.23916 (16) | 0.0440 (5) | |
| H3AA | 0.549 (3) | 0.657 (3) | 0.236 (2) | 0.067 (8)* | |
| C1B | 0.28710 (18) | 0.17323 (18) | 0.48905 (16) | 0.0351 (4) | |
| C2B | 0.1618 (2) | 0.3114 (2) | 0.5268 (2) | 0.0451 (5) | |
| H2B | 0.122 (2) | 0.382 (2) | 0.5441 (19) | 0.050 (7)* | |
| C3B | 0.2771 (2) | 0.2975 (2) | 0.50783 (18) | 0.0410 (5) | |
| H3B | 0.336 (3) | 0.354 (3) | 0.508 (2) | 0.061 (8)* | |
| N1B | 0.37852 (18) | 0.11030 (18) | 0.46300 (18) | 0.0472 (5) | |
| H1BA | 0.450 (3) | 0.147 (2) | 0.476 (2) | 0.053 (7)* | |
| H1BB | 0.375 (3) | 0.029 (3) | 0.472 (2) | 0.060 (8)* | |
| N2B | 0.18060 (15) | 0.11865 (16) | 0.49623 (14) | 0.0372 (4) | |
| H2BA | 0.164 (2) | 0.034 (3) | 0.4985 (19) | 0.058 (8)* | |
| N3B | 0.10420 (17) | 0.20371 (16) | 0.52047 (16) | 0.0431 (4) | |
| H3BA | 0.024 (3) | 0.184 (2) | 0.5222 (19) | 0.051 (7)* | |
| C4A | 0.58112 (19) | 0.40965 (18) | 0.26078 (18) | 0.0409 (5) | |
| C5A | 0.4574 (2) | 0.4142 (2) | 0.29303 (18) | 0.0422 (5) | |
| F1A1 | 0.4274 (7) | 0.3086 (7) | 0.3249 (5) | 0.0619 (13) | 0.639 (12) |
| F1A2 | 0.3945 (12) | 0.3073 (14) | 0.2818 (12) | 0.075 (3) | 0.361 (12) |
| F2A1 | 0.3589 (8) | 0.4515 (8) | 0.2173 (6) | 0.0681 (17) | 0.639 (12) |
| F2A2 | 0.3687 (14) | 0.4932 (10) | 0.2363 (13) | 0.067 (3) | 0.361 (12) |
| F3A1 | 0.4720 (7) | 0.4955 (5) | 0.3687 (6) | 0.0651 (13) | 0.639 (12) |
| F3A2 | 0.4800 (14) | 0.4446 (15) | 0.3885 (10) | 0.089 (4) | 0.361 (12) |
| O1A | 0.63921 (15) | 0.31128 (13) | 0.27322 (15) | 0.0526 (5) | |
| O2A | 0.60816 (16) | 0.50747 (14) | 0.22685 (17) | 0.0622 (5) | |
| C4B | 0.23375 (19) | 0.8115 (2) | 0.48329 (18) | 0.0413 (5) | |
| C5B | 0.2075 (2) | 0.6750 (2) | 0.45833 (19) | 0.0447 (5) | |
| F1B1 | 0.1329 (4) | 0.6255 (3) | 0.5045 (6) | 0.077 (2) | 0.639 (12) |
| F1B2 | 0.2028 (16) | 0.6103 (8) | 0.5367 (6) | 0.101 (4) | 0.361 (12) |
| F2B1 | 0.3162 (3) | 0.6097 (3) | 0.4866 (5) | 0.0669 (13) | 0.639 (12) |
| F2B2 | 0.2821 (9) | 0.6190 (6) | 0.4167 (12) | 0.089 (4) | 0.361 (12) |
| F3B1 | 0.1581 (7) | 0.6629 (3) | 0.3597 (3) | 0.090 (2) | 0.639 (12) |
| F3B2 | 0.0855 (7) | 0.6537 (6) | 0.3935 (8) | 0.079 (3) | 0.361 (12) |
| O1B | 0.13834 (14) | 0.86991 (13) | 0.49109 (15) | 0.0530 (5) | |
| O2B | 0.34134 (14) | 0.84860 (15) | 0.48973 (15) | 0.0538 (5) | |
| | | | | | |

Atomic displacement parameters $(Å^2)$

| | U^{11} | U^{22} | U^{33} | U^{12} | U^{13} | U^{23} |
|-----|-------------|-------------|-------------|-------------|-------------|--------------|
| C1A | 0.0374 (10) | 0.0299 (10) | 0.0508 (12) | 0.0050 (8) | 0.0177 (9) | 0.0021 (8) |
| C2A | 0.0330 (11) | 0.0431 (13) | 0.0629 (15) | -0.0022 (9) | 0.0194 (10) | 0.0043 (10) |
| C3A | 0.0351 (10) | 0.0382 (12) | 0.0595 (14) | 0.0072 (9) | 0.0208 (10) | 0.0022 (10) |
| N1A | 0.0434 (11) | 0.0275 (10) | 0.113 (2) | 0.0008 (8) | 0.0358 (12) | -0.0044 (10) |
| N2A | 0.0340 (9) | 0.0284 (9) | 0.0624 (12) | 0.0006 (7) | 0.0238 (8) | 0.0023 (8) |
| N3A | 0.0385 (9) | 0.0281 (9) | 0.0701 (13) | -0.0011 (7) | 0.0239 (9) | 0.0017 (8) |
| C1B | 0.0265 (9) | 0.0353 (11) | 0.0453 (11) | -0.0029 (7) | 0.0138 (8) | 0.0041 (8) |
| C2B | 0.0464 (12) | 0.0299 (11) | 0.0663 (15) | -0.0029 (9) | 0.0280 (11) | -0.0050 (10) |
| C3B | 0.0344 (10) | 0.0341 (11) | 0.0570 (13) | -0.0096 (8) | 0.0178 (9) | -0.0010 (9) |
| | | | | | | |

| N1B | 0.0302 (9) | 0.0350 (10) | 0.0823 (15) | -0.0018 (8) | 0.0259 (9) | 0.0024 (9) |
|------|-------------|-------------|-------------|--------------|-------------|--------------|
| N2B | 0.0288 (8) | 0.0279 (9) | 0.0588 (11) | -0.0013 (6) | 0.0192 (7) | -0.0009 (7) |
| N3B | 0.0345 (9) | 0.0330 (9) | 0.0709 (13) | -0.0024 (7) | 0.0298 (9) | -0.0039 (8) |
| C4A | 0.0354 (10) | 0.0301 (11) | 0.0639 (14) | -0.0021 (8) | 0.0252 (10) | -0.0036 (9) |
| C5A | 0.0363 (11) | 0.0397 (12) | 0.0549 (13) | -0.0014 (9) | 0.0201 (10) | 0.0010 (9) |
| F1A1 | 0.053 (3) | 0.0486 (17) | 0.099 (4) | -0.004 (2) | 0.045 (3) | 0.015 (3) |
| F1A2 | 0.046 (5) | 0.054 (3) | 0.135 (10) | -0.010 (4) | 0.044 (5) | 0.013 (7) |
| F2A1 | 0.0376 (17) | 0.094 (5) | 0.071 (2) | 0.011 (3) | 0.0149 (16) | 0.019 (3) |
| F2A2 | 0.038 (4) | 0.058 (5) | 0.113 (8) | 0.009 (4) | 0.035 (5) | 0.018 (4) |
| F3A1 | 0.061 (2) | 0.067 (3) | 0.079 (4) | -0.001 (2) | 0.039 (2) | -0.027 (2) |
| F3A2 | 0.070 (4) | 0.137 (11) | 0.069 (5) | -0.006 (7) | 0.034 (3) | -0.015 (7) |
| 01A | 0.0452 (9) | 0.0277 (8) | 0.0968 (13) | 0.0022 (6) | 0.0394 (9) | 0.0030 (7) |
| O2A | 0.0526 (10) | 0.0302 (8) | 0.1222 (16) | 0.0029 (7) | 0.0538 (11) | 0.0110 (9) |
| C4B | 0.0309 (10) | 0.0341 (11) | 0.0634 (14) | 0.0007 (8) | 0.0211 (10) | 0.0020 (9) |
| C5B | 0.0379 (11) | 0.0350 (12) | 0.0636 (15) | 0.0034 (9) | 0.0193 (10) | -0.0015 (10) |
| F1B1 | 0.071 (2) | 0.0309 (14) | 0.151 (6) | -0.0066 (15) | 0.068 (3) | 0.005 (2) |
| F1B2 | 0.174 (11) | 0.053 (4) | 0.069 (4) | -0.026 (6) | 0.025 (6) | 0.011 (3) |
| F2B1 | 0.0527 (15) | 0.0417 (14) | 0.107 (3) | 0.0162 (11) | 0.0249 (19) | -0.0071 (17) |
| F2B2 | 0.072 (6) | 0.059 (3) | 0.163 (11) | -0.002 (3) | 0.076 (7) | -0.038 (5) |
| F3B1 | 0.113 (5) | 0.0658 (19) | 0.069 (2) | -0.017 (2) | -0.003 (2) | -0.0142 (14) |
| F3B2 | 0.050 (3) | 0.060 (3) | 0.108 (6) | 0.002 (2) | -0.003 (3) | -0.033 (3) |
| O1B | 0.0343 (8) | 0.0290 (8) | 0.1059 (14) | -0.0020 (6) | 0.0366 (8) | -0.0059 (8) |
| O2B | 0.0310 (8) | 0.0434 (9) | 0.0937 (13) | -0.0013 (6) | 0.0290 (8) | 0.0035 (8) |
| | | | | | | |

Geometric parameters (Å, °)

| C1A—C3A | 1.399 (3) | N2B—H2BA | 0.95 (3) |
|----------|-----------|----------|------------|
| C1A—N1A | 1.344 (3) | N2B—N3B | 1.358 (2) |
| C1A—N2A | 1.338 (3) | N3B—H3BA | 0.91 (3) |
| C2A—H2A | 0.96 (3) | C4A—C5A | 1.547 (3) |
| C2A—C3A | 1.354 (3) | C4A—O1A | 1.234 (2) |
| C2A—N3A | 1.329 (3) | C4A—O2A | 1.238 (3) |
| СЗА—НЗА | 0.94 (3) | C5A—F1A1 | 1.312 (8) |
| N1A—H1AA | 0.85 (3) | C5A—F1A2 | 1.341 (14) |
| N1A—H1AB | 0.91 (3) | C5A—F2A1 | 1.309 (8) |
| N2A—H2AA | 0.94 (3) | C5A—F2A2 | 1.351 (15) |
| N2A—N3A | 1.357 (2) | C5A—F3A1 | 1.337 (7) |
| N3A—H3AA | 0.93 (3) | C5A—F3A2 | 1.298 (14) |
| C1B—C3B | 1.393 (3) | C4B—C5B | 1.538 (3) |
| C1B—N1B | 1.349 (3) | C4B—O1B | 1.255 (2) |
| C1B—N2B | 1.338 (2) | C4B—O2B | 1.221 (2) |
| C2B—H2B | 0.95 (3) | C5B—F1B1 | 1.295 (5) |
| C2B—C3B | 1.372 (3) | C5B—F1B2 | 1.299 (8) |
| C2B—N3B | 1.325 (3) | C5B—F2B1 | 1.337 (4) |
| СЗВ—НЗВ | 0.89 (3) | C5B—F2B2 | 1.284 (6) |
| N1B—H1BA | 0.84 (3) | C5B—F3B1 | 1.298 (4) |
| N1B—H1BB | 0.90 (3) | C5B—F3B2 | 1.375 (6) |
| | | | |

| N1A—C1A—C3A | 131.41 (19) | C2B—N3B—N2B | 108.01 (17) |
|-----------------------------------------|-------------------------|------------------------------------------------------|---------------------|
| N2A—C1A—C3A | 107.28 (18) | C2B—N3B—H3BA | 130.5 (16) |
| N2A—C1A—N1A | 121.30 (19) | N2B—N3B—H3BA | 121.1 (16) |
| C3A—C2A—H2A | 129.0 (15) | O1A—C4A—C5A | 116.52 (18) |
| N3A—C2A—H2A | 121.0 (15) | O1A—C4A—O2A | 129.27 (19) |
| N3A—C2A—C3A | 109.9 (2) | O2A—C4A—C5A | 114.21 (17) |
| С1А—С3А—Н3А | 127.7 (15) | F1A1—C5A—C4A | 113.4 (4) |
| C2A - C3A - C1A | 106.02 (19) | F1A1—C5A—F3A1 | 108.0 (4) |
| $C_2A - C_3A - H_3A$ | 126.3 (15) | F1A2-C5A-C4A | 113.7(7) |
| C1A $N1A$ $H1AA$ | 120.3(19) 118 4 (19) | F1A2 $C5A$ $F2A2$ | 103.9(7) |
| C14 N14 H14R | 110.1 (19) | $F_{2A1} C_{5A} C_{4A}$ | 103.9(7) |
| $H1\Delta\Delta$ $N1\Delta$ $H1\DeltaB$ | 120(3) | F_{2A1} C_{5A} F_{1A1} | 108.0(4) |
| C_{1A} N2A H2AA | 120(3) 128.8(18) | $F_{2A1} = C_{5A} = F_{2A1}$ | 106.0(4) |
| C1A = N2A = N2A | 120.0(10) | $F_2A_1 = C_5A = F_5A_1$ | 100.2(4) |
| NIA NIA HIAA | 109.00(17) | F2A2 - C3A - C4A | 113.1(7) |
| $N_{3}A - N_{2}A - H_{2}AA$ | 122.2(18) | $F_{3A1} = C_{3A} = C_{4A}$ | 109.7(3) |
| C_{2A} N3A N2A | 107.79 (18) | F3A2 - C3A - C4A | 112.6 (6) |
| C2A—N3A—H3AA | 129.0 (18) | F3A2—C5A—F1A2 | 105.7 (7) |
| N2A—N3A—H3AA | 122.6 (18) | F3A2—C5A—F2A2 | 107.3 (8) |
| N1B—C1B—C3B | 130.75 (19) | O1B—C4B—C5B | 114.19 (18) |
| N2B—C1B—C3B | 107.58 (17) | O2B—C4B—C5B | 116.60 (19) |
| N2B—C1B—N1B | 121.60 (19) | O2B—C4B—O1B | 129.2 (2) |
| C3B—C2B—H2B | 131.0 (15) | F1B1—C5B—C4B | 113.5 (3) |
| N3B—C2B—H2B | 119.5 (15) | F1B1—C5B—F2B1 | 105.8 (3) |
| N3B—C2B—C3B | 109.57 (19) | F1B1—C5B—F3B1 | 110.1 (3) |
| C1B—C3B—H3B | 125.5 (18) | F1B2—C5B—C4B | 113.3 (4) |
| C2B—C3B—C1B | 105.75 (18) | F1B2—C5B—F3B2 | 99.4 (5) |
| C2B—C3B—H3B | 128.7 (18) | F2B1—C5B—C4B | 111.4 (2) |
| C1B—N1B—H1BA | 114.8 (18) | F2B2—C5B—C4B | 117.5 (3) |
| C1B—N1B—H1BB | 113.5 (17) | F2B2—C5B—F1B2 | 107.4 (6) |
| H1BA—N1B—H1BB | 121 (2) | F2B2—C5B—F3B2 | 104.7 (5) |
| C1B—N2B—H2BA | 128.4 (16) | F3B1—C5B—C4B | 108.6 (2) |
| C1B—N2B—N3B | 109.08 (17) | F3B1—C5B—F2B1 | 107.2 (3) |
| N3B—N2B—H2BA | 121.5 (16) | F3B2—C5B—C4B | 112.7(3) |
| | 121.0 (10) | | 112.7 (3) |
| C1A—N2A—N3A—C2A | -10(3) | O1A - C4A - C5A - F3A2 | 88 7 (8) |
| C_{3A} C_{1A} N_{2A} N_{3A} | 0.6(3) | $O^2A - C^4A - C^5A - F^1A^1$ | 1774(4) |
| $C_{3}A = C_{2}A = N_{3}A = N_{2}A$ | 11(3) | O2A = C4A = C5A = F1A2 | 1/7.4(4) 1494(7) |
| $N_{1A} = C_{2A} = N_{3A} = N_{2A}$ | 1.1(3) 178 8 (3) | $O_{2A} = C_{4A} = C_{5A} = 11A_2$ | 55.5(5) |
| N1A C1A N2A N3A | -1783(2) | $O_{2A} = C_{4A} = C_{5A} = \Gamma_{2A} \Gamma_{2A}$ | 31.3(7) |
| NIA = CIA = NZA = NJA | 170.3(2) | $O_{2A} = C_{4A} = C_{5A} = \Gamma_{2A2}$ | -617(4) |
| NZA = CIA = CJA = CIA | 0.1(3) | $O_{2A} = C_{4A} = C_{5A} = F_{5A1}$ | -01.7(4) |
| $N_{3A} = C_{2A} = C_{3A} = C_{1A}$ | -0.7(3) | O_{2A} C_{4A} C_{5A} F_{5A2} | -90.3(8) |
| CIB = N2B = N3B = C2B | 1.0(3) | OIB - C4B - C5B - FIBI | -38.2 (4) |
| C_{3B} C_{1B} N_{2B} N_{3B} | -0.9(2) | 01B-04B-05B-F1B2 | -/6.1 (9) |
| C3B—C2B—N3B—N2B | -0.6 (3) | 01B | -157.5 (3) |
| NIB—CIB—C3B—C2B | 177.5 (2) | 01B—C4B—C5B—F2B2 | 157.7 (8) |
| N1B—C1B—N2B—N3B | -178.2(2) | O1B—C4B—C5B—F3B1 | 84.6 (5) |
| N2B—C1B—C3B—C2B | 0.5 (3) | O1B—C4B—C5B—F3B2 | 35.9 (7) |
| N3B—C2B—C3B—C1B | 0.1 (3) | O2B—C4B—C5B—F1B1 | 143.9 (4) |

supporting information

| O1A—C4A—C5A—F1A1 | -3.4(4) | O2B—C4B—C5B—F1B2 | 106.0 (9) |
|------------------|------------|----------------------------------------------------------|------------|
| O1A—C4A—C5A—F1A2 | -31.4(7) | O2B—C4B—C5B—F2B1 | 24.5 (4) |
| 01A—C4A—C5A—F2A1 | -125.3 (4) | O2B C4B C5B F2B1 O2B C4B C5B F2B2 O2B C4B C5B F2B1 | -20.3(9) |
| 01A—C4A—C5A—F2A2 | -149.5 (6) | O2B—C4B—C5B—F3B1 | -93.3 (5) |
| 01A—C4A—C5A—F3A1 | 117.5 (4) | O2B—C4B—C5B—F3B2 | -142.1 (6) |

Hydrogen-bond geometry (Å, °)

| <i>D</i> —H··· <i>A</i> | D—H | Н…А | D···A | <i>D</i> —H··· <i>A</i> |
|------------------------------------------|----------|----------|-----------|-------------------------|
| N1A—H1AA····O1A ⁱ | 0.85 (3) | 2.28 (3) | 2.936 (3) | 134 (2) |
| $N1A$ — $H1AB$ ···O2 A^{ii} | 0.91 (3) | 1.99 (3) | 2.884 (3) | 169 (3) |
| $N2A$ — $H2AA$ ···O1 A^{ii} | 0.94 (3) | 1.85 (3) | 2.778 (2) | 171 (3) |
| N3 <i>A</i> —H3 <i>AA</i> ···O2 <i>A</i> | 0.93 (3) | 1.78 (3) | 2.705 (2) | 172 (3) |
| N1 B —H1 BA ···O2 B ⁱⁱⁱ | 0.84 (3) | 2.18 (3) | 2.962 (2) | 153 (2) |
| N1B—H1BB····O2B ^{iv} | 0.90 (3) | 2.03 (3) | 2.929 (3) | 173 (2) |
| N2B—H2BA····O1B ^{iv} | 0.95 (3) | 1.81 (3) | 2.756 (2) | 174 (2) |
| $N3B$ — $H3BA$ ···O1 B^{v} | 0.91 (3) | 1.82 (3) | 2.728 (2) | 171 (2) |

Symmetry codes: (i) x, y+1, z; (ii) -x+3/2, y+1/2, -z+1/2; (iii) -x+1, -y+1, -z+1; (iv) x, y-1, z; (v) -x, -y+1, -z+1.