organic compounds

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1-Methylpiperazine-1,4-diium dipicrate

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Key indicators: single-crystal X-ray study; T = 295 K; mean σ (C–C) = 0.002 Å; R factor = 0.044; wR factor = 0.123; data-to-parameter ratio = 11.5.

In the crystal structure of the title compound [systematic name: 1-methylpiperazine-1,4-diium bis(2,4,6-trinitrophenolate)], $C_5H_{14}N_2^{2+} \cdot 2C_6H_2N_3O_7^-$, the ionic components are connected by relatively strong N-H···O hydrogen bonds into centrosymmetric six-membered conglomerates, which comprise two dications and four anions. Besides Coulombic interactions, only weak C-H···O interactions and some stacking between picrates (separation between the planes of ca. 3.4 Å but only a small overlapping) can be identified between these 'building blocks' of the crystal structure. The piperazine ring adopts a chair conformation with the methyl substituent in the equatorial position. In the picrate anions, the twist angles of the nitro groups depend on their positions relative to the phenolate O atom: it is much smaller for the NO₂ groups *para* to the C $-O^-$ group [15.23 (9) and 3.92 (14)°] than for the groups in the ortho positions [28.76 (13)-39.84 (11)°].

Related literature

For examples of the biological activity of piperazines: Brockunier *et al.* (2004); Bogatcheva *et al.* (2006). For the crystal structures of simple piperidinium picrates, see: Fun *et al.* (2010); Li *et al.* (2009); Verdonk *et al.* (1997); Wang & Jia (2008). For a description of the Cambridge Structural Database, see: Allen (2002). For asymmetry parameters, see: Duax & Norton (1975).



 $\gamma = 81.558 \ (12)^{\circ}$

Z = 2

V = 1109.6 (3) Å³

Mo $K\alpha$ radiation

 $0.4 \times 0.15 \times 0.07 \text{ mm}$

21056 measured reflections

4891 independent reflections

3624 reflections with $I > 2\sigma(I)$

 $\mu = 0.15 \text{ mm}^{-3}$ T = 295 K

 $R_{\rm int} = 0.021$

Experimental

Crvstal data

 $C_{5}H_{14}N_{2}^{2+}\cdot 2C_{6}H_{2}N_{3}O_{7}^{-}$ $M_{r} = 558.39$ Triclinic, $P\overline{1}$ a = 8.2001 (12) Å b = 10.1780 (15) Å c = 13.7399 (18) Å a = 89.798 (12)° $\beta = 78.130$ (11)°

Data collection

Oxford Diffraction Xcalibur Eos diffractometer Absorption correction: multi-scan (*CrysAlis PRO*; Oxford Diffraction, 2009)

 $T_{\min} = 0.936, \ T_{\max} = 1.000$

Refinement

| $R[F^2 > 2\sigma(F^2)] = 0.044$ | H atoms treated by a mixture of |
|---------------------------------|--|
| $wR(F^2) = 0.123$ | independent and constrained |
| S = 0.95 | refinement |
| 4891 reflections | $\Delta \rho_{\rm max} = 0.24 \text{ e } \text{\AA}^{-3}$ |
| 424 parameters | $\Delta \rho_{\rm min} = -0.30 \ {\rm e} \ {\rm \AA}^{-3}$ |

Table 1

Hydrogen-bond geometry (Å, °).

| $D - H \cdots A$ | D-H | $H \cdots A$ | $D \cdots A$ | $D - \mathbf{H} \cdot \cdot \cdot A$ |
|---------------------------------|------------|--------------|--------------|--------------------------------------|
| N11-H11···O1A | 0.892 (18) | 1.831 (18) | 2.6305 (17) | 148.1 (16) |
| $N11 - H11 \cdots O22A$ | 0.892 (18) | 2.356 (18) | 2.996 (2) | 128.8 (14) |
| $N14-H14B\cdotsO1B^{i}$ | 0.88 (2) | 1.98 (2) | 2.8181 (19) | 157.3 (17) |
| $N14-H14A\cdotsO1B$ | 0.92 (2) | 1.99 (2) | 2.7962 (18) | 146.4 (18) |
| N14 $-$ H14 A ···O22 B | 0.92 (2) | 2.28 (2) | 2.992 (2) | 133.9 (16) |
| $C5A - H5A \cdots O21A^{ii}$ | 0.917 (19) | 2.476 (19) | 3.383 (2) | 170.3 (16) |
| $C5B - H5B \cdots O21B^{iii}$ | 0.913 (18) | 2.487 (18) | 3.394 (2) | 172.3 (15) |
| $C11A - H11C \cdots O41A^{iv}$ | 0.93 (3) | 2.48 (3) | 3.345 (2) | 155 (2) |
| $C11A - H11A \cdots O62A^{iii}$ | 0.94 (3) | 2.57 (3) | 3.496 (3) | 168 (2) |
| $C13-H13A\cdots O62B^{v}$ | 0.96 (2) | 2.46 (2) | 3.386 (2) | 162.9 (17) |

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

Data collection: *CrysAlis PRO* (Oxford Diffraction, 2009); cell refinement: *CrysAlis PRO*; data reduction: *CrysAlis PRO*; program(s) used to solve structure: *SIR92* (Altomare *et al.*, 1993); program(s) used to refine structure: *SHELXL97* (Sheldrick, 2008); molecular graphics: *Stereochemical Workstation Operation Manual* (Siemens, 1989); software used to prepare material for publication: *SHELXL97*.

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Supplementary data and figures for this paper are available from the IUCr electronic archives (Reference: FL2330).

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1-Methylpiperazine-1,4-diium dipicrate

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S1. Comment

Piperazines are among the most important building blocks in today's drug discovery. They are found in biologically active compounds across a number of different therapeutic areas such as antifungal, antibacterial, antimalarial, antipsychotic, antidepressant and antitumour activity against colon, prostate, breast, lung and leukemia tumors (for instance, Brockunier *et al.*, 2004, Bogatcheva *et al.*, 2006). A small number of piperazinium picrates or piperazinediium dipicrates have been structurally characterized, however generally the cations were heavily substituted. On the other hand, picric acid ($pK_a=0.38$) has been studied for its ability to form salts which display wide spectrum of intermolecular interactions, for instance hydrogen bonds of different strengths and/or $\pi \cdots \pi$ stacking interactions. In the course of our studies of picrates of simple organic cations we have determined the crystal and molecular structure of the title compound (I: 1-methyl-piprazinediium di(2,4,6-trinitrophenolate), Scheme 1).

In the CSD (Allen, 2002; Version 5.31 of Nov. 2009, updated August 2010) there are only a few picrates of simple piperazinium derivatives, for instance 4-(4-carboxybenzyl)-1-methylpiperazin-1-ium picrate (Li *et al.*, 2009), 1-(2-meth-oxyphenyl)piperazinium picrate (Verdonk *et al.*, 1997) or piperazine-1,4-diium–dipicrate piperazine complex (Wang & Jia, 2008). Also some more complicated structures were reported, for instance 4-(3-Carboxy-1-ethyl-6-fluoro-4-oxo-1,4-dihydro-7-quinolyl)-1-methylpiperazinium picrate (Fun *et al.*, 2010).

In the crystal structure **I** there are two picrate anions and 1-methylpiperidinediium dication (Fig. 1); the presence of ionic species is supported by the successful location and refinement of the hydrogen atoms at both nitrogen atoms in the piperidine ring as well as by inspection of the pattern of bond distances and angles. The piperazine ring adopts an almost ideal chair conformation; the values of asymmetry parameters (Duax & Norton, 1975), which measure the deviations from the ideal symmetry (in the case $D_{3 d}$), are very small, less than 1.6°. The methyl substituent is in the equatorial position as can be seen from the torsion angles C13—C12—C11—C11A: 176.60 (15)° and C15—C16—C11—C11A: -176.72 (14)°. Both aromatic rings are in a good approximation planar, maximum deviation from the least-squares plane calculated by the six ring atoms is 0.0248 (11)Å in the anion A and 0.0297 (10)Å in anion B. The nitro groups are twisted with respect to the ring planes, for the groups *ortho* with respect to the C—O[•] group (at C2 and C6) this twist is of course significantly larger (ranging from 28.76 (13)° to 39.84 (11)°) than for the groups in *para* positions, at C4 (15.23 (9)° in anion A, only 3.92 (14)° in B).

In the crystal structure the building block is made up of a centrosymetric pair of hydrogen bonded ionic components: two dications and four anions (Table 1, Fig. 2). Using graph set notation one can identify - taking into account the primary interactions only - the centrosymmetric ring $R^2_4(8)$ and dimeric D motifs. Interestingly no strong hydrogen bonds are observed between these structures; besides the coulombic interactions only weak C—H…O and some stacking between picrates (Fig. 3) organize the crystal packing.

S2. Experimental

1-Methyl piperazine (1.00 g, 0.01 mol) was dissolved in 20 ml of alcohol. Picric acid (4.58 g, 0.02 mol) was dissolved in 50 ml of water. Both the solutions were mixed and to this, 5 ml of 3M HCl was added and stirred for few minutes. The formed complex was filtered and dried, crystals appropriate for X-ray data collection were found without further recrystallization (m. p. >523 K). Composition: Found (Calculated): C: 36.48 (36.57); H: 3.20 (3.25); N:19.98 (20.07).

S3. Refinement

Hydrogen atoms were located in difference Fourier maps and isotropically refined.



Figure 1

Anisotropic ellipsoid representation of the ionic components of I together with atom labelling scheme. The ellipsoids are drawn at 50% probability level, hydrogen atoms are depicted as spheres with arbitrary radii; hydrogen bonds are shown as dashed lines.



Figure 2

The centrosymmetric dimer of salt I; hydrogen bonds are shown as dashed lines. Symmetry codes: (i) -x, 1 - y, 1 - z.



Z = 2

F(000) = 576

 $\theta = 3.0 - 28.0^{\circ}$

 $\mu = 0.15 \text{ mm}^{-1}$

Block, yellow

 $0.4 \times 0.15 \times 0.07 \text{ mm}$

T = 295 K

 $D_{\rm x} = 1.671 {\rm Mg m^{-3}}$

Mo *K* α radiation, $\lambda = 0.71073$ Å

Cell parameters from 12041 reflections

Figure 3

The crystal packing as seen approximately along y-direction. Hydrogen bonds are shown as dashed lines.

1-methylpiperazine-1,4-diium bis(2,4,6-trinitrophenolate)

Crystal data

 $C_{5}H_{14}N_{2}^{2+} \cdot 2C_{6}H_{2}N_{3}O_{7}^{-}$ $M_{r} = 558.39$ Triclinic, *P*1 Hall symbol: -P 1 a = 8.2001 (12) Å b = 10.1780 (15) Å c = 13.7399 (18) Å $a = 89.798 (12)^{\circ}$ $\beta = 78.130 (11)^{\circ}$ $\gamma = 81.558 (12)^{\circ}$ $V = 1109.6 (3) \text{ Å}^{3}$

Data collection

| Oxford Diffraction Xcalibur Eos | 21056 measured reflections |
|--|---|
| diffractometer | 4891 independent reflections |
| Radiation source: Enhance (Mo) X-ray Source | 3624 reflections with $I > 2\sigma(I)$ |
| Graphite monochromator | $R_{\rm int} = 0.021$ |
| Detector resolution: 16.1544 pixels mm ⁻¹ | $\theta_{\rm max} = 28.0^{\circ}, \ \theta_{\rm min} = 3.0^{\circ}$ |
| ω scans | $h = -10 \rightarrow 10$ |
| Absorption correction: multi-scan | $k = -13 \rightarrow 12$ |
| (CrysAlis PRO; Oxford Diffraction, 2009) | $l = -18 \rightarrow 18$ |
| $T_{\min} = 0.936, \ T_{\max} = 1.000$ | |

Refinement

Refinement on F^2 Least-squares matrix: full $R[F^2 > 2\sigma(F^2)] = 0.044$ $wR(F^2) = 0.123$ S = 0.954891 reflections 424 parameters 0 restraints Primary atom site location: structure-invariant direct methods Secondary atom site location: difference Fourier map Hydrogen site location: inferred from neighbouring sites H atoms treated by a mixture of independent and constrained refinement $w = 1/[\sigma^2(F_o^2) + (0.0725P)^2 + 0.3607P]$ where $P = (F_o^2 + 2F_c^2)/3$ $(\Delta/\sigma)_{max} < 0.001$ $\Delta\rho_{max} = 0.24$ e Å⁻³ $\Delta\rho_{min} = -0.30$ e Å⁻³

Special details

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

Refinement. Refinement of F^2 against ALL reflections. The weighted *R*-factor *wR* and goodness of fit *S* are based on F^2 , conventional *R*-factors *R* are based on *F*, with *F* set to zero for negative F^2 . The threshold expression of $F^2 > \sigma(F^2)$ is used only for calculating *R*-factors(gt) *etc.* and is not relevant to the choice of reflections for refinement. *R*-factors based on F^2 are statistically about twice as large as those based on *F*, and *R*- factors based on ALL data will be even larger.

| | x | у | Ζ | $U_{ m iso}$ */ $U_{ m eq}$ | |
|------|---------------|---------------|--------------|-----------------------------|--|
| C1A | -0.10838 (19) | 0.86643 (15) | 0.10686 (11) | 0.0303 (3) | |
| O1A | -0.02003 (16) | 0.75573 (12) | 0.08429 (10) | 0.0469 (3) | |
| C2A | -0.04660 (18) | 0.98367 (16) | 0.13439 (12) | 0.0309 (3) | |
| N2A | 0.12392 (16) | 0.97199 (15) | 0.15030 (11) | 0.0394 (3) | |
| O21A | 0.19522 (16) | 1.07010 (15) | 0.13867 (13) | 0.0601 (4) | |
| O22A | 0.18805 (16) | 0.86647 (14) | 0.17951 (12) | 0.0568 (4) | |
| C3A | -0.1404 (2) | 1.10787 (16) | 0.14843 (12) | 0.0314 (3) | |
| H3A | -0.091 (3) | 1.179 (2) | 0.1651 (16) | 0.052 (6)* | |
| C4A | -0.30793 (18) | 1.12289 (14) | 0.14285 (11) | 0.0284 (3) | |
| N4A | -0.40990 (18) | 1.25160 (13) | 0.16554 (10) | 0.0350 (3) | |
| O41A | -0.56396 (15) | 1.25745 (13) | 0.18252 (10) | 0.0474 (3) | |
| O42A | -0.33873 (18) | 1.34906 (12) | 0.16819 (12) | 0.0566 (4) | |
| C5A | -0.38244 (19) | 1.01553 (15) | 0.12063 (11) | 0.0288 (3) | |
| H5A | -0.495 (2) | 1.0237 (18) | 0.1186 (13) | 0.036 (5)* | |
| C6A | -0.28521 (19) | 0.89319 (14) | 0.10388 (11) | 0.0294 (3) | |
| N6A | -0.36772 (18) | 0.78285 (13) | 0.08081 (11) | 0.0370 (3) | |
| O61A | -0.3268 (2) | 0.67315 (13) | 0.11180 (13) | 0.0662 (5) | |
| O62A | -0.47823 (18) | 0.80553 (14) | 0.03359 (11) | 0.0557 (4) | |
| C1B | 0.29112 (18) | 0.24867 (15) | 0.41703 (11) | 0.0270 (3) | |
| O1B | 0.19899 (13) | 0.35763 (11) | 0.44572 (8) | 0.0353 (3) | |
| C2B | 0.23173 (18) | 0.13264 (16) | 0.38650 (12) | 0.0299 (3) | |
| N2B | 0.05928 (16) | 0.14283 (15) | 0.37308 (12) | 0.0404 (3) | |
| O21B | -0.00902 (16) | 0.04406 (15) | 0.38274 (15) | 0.0690 (5) | |
| O22B | -0.00859 (16) | 0.24821 (14) | 0.34659 (12) | 0.0576 (4) | |
| C3B | 0.32794 (19) | 0.00995 (16) | 0.36639 (12) | 0.0315 (3) | |
| H3B | 0.279 (2) | -0.0637 (19) | 0.3481 (14) | 0.040 (5)* | |
| C4B | 0.49633 (19) | -0.00422 (15) | 0.37139 (11) | 0.0301 (3) | |
| N4B | 0.59846 (18) | -0.13327 (14) | 0.34997 (11) | 0.0389 (3) | |
| O41B | 0.74541 (16) | -0.14586 (14) | 0.35957 (12) | 0.0558 (4) | |
| O42B | 0.53534 (19) | -0.22459 (14) | 0.32327 (14) | 0.0652 (4) | |
| C5B | 0.56874 (19) | 0.10308 (16) | 0.39575 (11) | 0.0303 (3) | |
| H5B | 0.681 (2) | 0.0949 (17) | 0.3956 (13) | 0.033 (4)* | |
| C6B | 0.46974 (18) | 0.22393 (16) | 0.41531 (11) | 0.0295 (3) | |
| N6B | 0.55497 (17) | 0.33587 (15) | 0.43280 (12) | 0.0414 (4) | |
| O61B | 0.5170(2) | 0.44149 (14) | 0.39533 (12) | 0.0590 (4) | |

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

| O62B | 0.66595 (18) | 0.31520 (16) | 0.47993 (14) | 0.0699 (5) |
|------|---------------|--------------|--------------|------------|
| N11 | 0.16733 (15) | 0.57565 (13) | 0.16779 (10) | 0.0291 (3) |
| H11 | 0.114 (2) | 0.6554 (18) | 0.1567 (13) | 0.030 (4)* |
| C11A | 0.2986 (3) | 0.5363 (2) | 0.07575 (15) | 0.0456 (5) |
| H11C | 0.346 (3) | 0.450 (3) | 0.0848 (19) | 0.071 (7)* |
| H11B | 0.243 (3) | 0.539 (2) | 0.024 (2) | 0.068 (7)* |
| H11A | 0.372 (3) | 0.600 (3) | 0.068 (2) | 0.079 (8)* |
| C12 | 0.2433 (2) | 0.58588 (17) | 0.25660 (13) | 0.0348 (4) |
| H12B | 0.300(2) | 0.498 (2) | 0.2664 (14) | 0.041 (5)* |
| H12A | 0.322 (3) | 0.647 (2) | 0.2416 (16) | 0.053 (6)* |
| C13 | 0.1088 (2) | 0.63356 (18) | 0.34660 (13) | 0.0387 (4) |
| H13B | 0.054 (2) | 0.7194 (19) | 0.3366 (13) | 0.035 (5)* |
| H13A | 0.157 (2) | 0.636 (2) | 0.4042 (16) | 0.049 (5)* |
| N14 | -0.02066 (18) | 0.54247 (15) | 0.36518 (11) | 0.0361 (3) |
| H14B | -0.100 (3) | 0.576 (2) | 0.4161 (16) | 0.044 (5)* |
| H14A | 0.031 (3) | 0.460 (2) | 0.3783 (15) | 0.048 (5)* |
| C15 | -0.0961 (2) | 0.52998 (19) | 0.27659 (13) | 0.0364 (4) |
| H15B | -0.153 (2) | 0.616 (2) | 0.2644 (15) | 0.042 (5)* |
| H15A | -0.171 (3) | 0.471 (2) | 0.2906 (15) | 0.047 (5)* |
| C16 | 0.0398 (2) | 0.48330 (17) | 0.18685 (13) | 0.0338 (3) |
| H16B | 0.098 (2) | 0.3954 (19) | 0.1948 (14) | 0.036 (5)* |
| H16A | -0.012 (3) | 0.481 (2) | 0.1283 (17) | 0.056 (6)* |
| | | | | |

Atomic displacement parameters $(Å^2)$

| | U^{11} | U^{22} | U^{33} | U^{12} | U^{13} | U^{23} |
|------|-------------|------------|-------------|-------------|--------------|-------------|
| C1A | 0.0339 (8) | 0.0278 (8) | 0.0286 (8) | 0.0045 (6) | -0.0116 (6) | 0.0010 (6) |
| O1A | 0.0507 (7) | 0.0349 (6) | 0.0543 (8) | 0.0144 (5) | -0.0239 (6) | -0.0092 (6) |
| C2A | 0.0257 (7) | 0.0361 (8) | 0.0310 (8) | -0.0019 (6) | -0.0081 (6) | 0.0030 (6) |
| N2A | 0.0275 (7) | 0.0450 (8) | 0.0462 (9) | -0.0034 (6) | -0.0097 (6) | -0.0010(7) |
| O21A | 0.0355 (7) | 0.0566 (9) | 0.0925 (12) | -0.0161 (6) | -0.0166 (7) | 0.0067 (8) |
| O22A | 0.0422 (7) | 0.0511 (8) | 0.0820 (11) | 0.0035 (6) | -0.0320 (7) | 0.0070 (7) |
| C3A | 0.0342 (8) | 0.0290 (8) | 0.0324 (8) | -0.0068 (6) | -0.0088 (6) | 0.0024 (6) |
| C4A | 0.0311 (7) | 0.0244 (7) | 0.0284 (8) | 0.0010 (6) | -0.0072 (6) | 0.0014 (6) |
| N4A | 0.0435 (8) | 0.0274 (7) | 0.0324 (7) | 0.0029 (6) | -0.0099 (6) | -0.0003 (5) |
| O41A | 0.0381 (7) | 0.0415 (7) | 0.0571 (8) | 0.0112 (5) | -0.0094 (6) | -0.0042 (6) |
| O42A | 0.0646 (9) | 0.0267 (6) | 0.0792 (11) | -0.0052 (6) | -0.0174 (8) | -0.0055 (6) |
| C5A | 0.0280 (7) | 0.0307 (8) | 0.0287 (8) | -0.0012 (6) | -0.0102 (6) | 0.0027 (6) |
| C6A | 0.0357 (8) | 0.0260 (7) | 0.0289 (8) | -0.0037 (6) | -0.0130 (6) | 0.0011 (6) |
| N6A | 0.0453 (8) | 0.0307 (7) | 0.0387 (8) | -0.0068 (6) | -0.0163 (6) | -0.0020 (6) |
| O61A | 0.0923 (11) | 0.0293 (7) | 0.0920 (12) | -0.0132 (7) | -0.0508 (10) | 0.0087 (7) |
| O62A | 0.0606 (8) | 0.0531 (8) | 0.0682 (9) | -0.0181 (7) | -0.0407 (8) | 0.0056 (7) |
| C1B | 0.0256 (7) | 0.0315 (8) | 0.0221 (7) | 0.0001 (6) | -0.0036 (5) | -0.0009 (6) |
| O1B | 0.0338 (6) | 0.0350 (6) | 0.0336 (6) | 0.0054 (5) | -0.0061 (5) | -0.0059 (5) |
| C2B | 0.0227 (7) | 0.0359 (8) | 0.0308 (8) | -0.0032 (6) | -0.0055 (6) | 0.0006 (6) |
| N2B | 0.0265 (7) | 0.0434 (8) | 0.0523 (9) | -0.0043 (6) | -0.0108 (6) | -0.0032 (7) |
| O21B | 0.0357 (7) | 0.0529 (9) | 0.1243 (15) | -0.0158 (6) | -0.0236 (8) | 0.0045 (9) |
| O22B | 0.0400 (7) | 0.0510 (8) | 0.0875 (11) | 0.0010 (6) | -0.0315 (7) | 0.0076 (7) |
| | | | | | | |

| C3B | 0.0312 (8) | 0.0310 (8) | 0.0331 (8) | -0.0061 (6) | -0.0070 (6) | -0.0003 (6) |
|------|-------------|-------------|-------------|-------------|-------------|-------------|
| C4B | 0.0288 (7) | 0.0312 (8) | 0.0278 (8) | 0.0027 (6) | -0.0050 (6) | -0.0014 (6) |
| N4B | 0.0388 (8) | 0.0355 (8) | 0.0370 (8) | 0.0050 (6) | -0.0031 (6) | -0.0023 (6) |
| O41B | 0.0365 (7) | 0.0511 (8) | 0.0751 (10) | 0.0147 (6) | -0.0155 (6) | -0.0074 (7) |
| O42B | 0.0597 (9) | 0.0352 (7) | 0.0992 (13) | 0.0017 (6) | -0.0186 (8) | -0.0210 (8) |
| C5B | 0.0227 (7) | 0.0402 (9) | 0.0264 (8) | 0.0003 (6) | -0.0053 (6) | -0.0026 (6) |
| C6B | 0.0273 (7) | 0.0349 (8) | 0.0264 (7) | -0.0051 (6) | -0.0056 (6) | -0.0037 (6) |
| N6B | 0.0318 (7) | 0.0453 (9) | 0.0462 (9) | -0.0065 (6) | -0.0052 (6) | -0.0166 (7) |
| O61B | 0.0718 (9) | 0.0448 (8) | 0.0644 (9) | -0.0235 (7) | -0.0133 (8) | 0.0015 (7) |
| O62B | 0.0495 (8) | 0.0688 (10) | 0.0995 (13) | 0.0001 (7) | -0.0399 (9) | -0.0343 (9) |
| N11 | 0.0286 (6) | 0.0248 (6) | 0.0310 (7) | 0.0034 (5) | -0.0047 (5) | -0.0002 (5) |
| C11A | 0.0464 (10) | 0.0415 (11) | 0.0383 (10) | 0.0077 (9) | 0.0055 (8) | 0.0011 (8) |
| C12 | 0.0280 (8) | 0.0350 (9) | 0.0423 (9) | -0.0012 (7) | -0.0122 (7) | -0.0009 (7) |
| C13 | 0.0431 (9) | 0.0357 (9) | 0.0386 (9) | 0.0020 (7) | -0.0169 (8) | -0.0092 (7) |
| N14 | 0.0343 (7) | 0.0384 (8) | 0.0289 (7) | 0.0082 (6) | -0.0010 (6) | -0.0024 (6) |
| C15 | 0.0271 (8) | 0.0411 (9) | 0.0403 (9) | -0.0024 (7) | -0.0073 (7) | 0.0020 (7) |
| C16 | 0.0376 (8) | 0.0309 (8) | 0.0342 (9) | -0.0054 (7) | -0.0104 (7) | -0.0033 (7) |
| | | | | | | |

Geometric parameters (Å, °)

| C1A—O1A | 1.2494 (18) | N4B—O42B | 1.220 (2) |
|----------|-------------|-----------|-------------|
| C1A—C2A | 1.443 (2) | N4B—O41B | 1.2271 (19) |
| C1A—C6A | 1.445 (2) | C5B—C6B | 1.365 (2) |
| C2A—C3A | 1.372 (2) | C5B—H5B | 0.913 (18) |
| C2A—N2A | 1.4470 (19) | C6B—N6B | 1.466 (2) |
| N2A—O21A | 1.2233 (19) | N6B—O62B | 1.215 (2) |
| N2A—O22A | 1.2283 (19) | N6B—O61B | 1.217 (2) |
| C3A—C4A | 1.378 (2) | N11—C16 | 1.490 (2) |
| СЗА—НЗА | 0.93 (2) | N11—C12 | 1.490 (2) |
| C4A—C5A | 1.391 (2) | N11—C11A | 1.495 (2) |
| C4A—N4A | 1.4445 (19) | N11—H11 | 0.892 (18) |
| N4A—O42A | 1.2267 (19) | C11A—H11C | 0.93 (3) |
| N4A—O41A | 1.2291 (18) | C11A—H11B | 0.92 (3) |
| C5A—C6A | 1.369 (2) | C11A—H11A | 0.94 (3) |
| C5A—H5A | 0.917 (19) | C12—C13 | 1.505 (2) |
| C6A—N6A | 1.4605 (19) | C12—H12B | 0.97 (2) |
| N6A—O62A | 1.2144 (18) | C12—H12A | 0.96 (2) |
| N6A—O61A | 1.2190 (19) | C13—N14 | 1.493 (2) |
| C1B—O1B | 1.2612 (18) | С13—Н13В | 0.946 (19) |
| C1B—C2B | 1.437 (2) | C13—H13A | 0.96 (2) |
| C1B—C6B | 1.445 (2) | N14—C15 | 1.488 (2) |
| C2B—C3B | 1.372 (2) | N14—H14B | 0.88 (2) |
| C2B—N2B | 1.4521 (19) | N14—H14A | 0.92 (2) |
| N2B—O21B | 1.215 (2) | C15—C16 | 1.507 (2) |
| N2B—O22B | 1.2238 (19) | C15—H15B | 0.96 (2) |
| C3B—C4B | 1.383 (2) | C15—H15A | 0.92 (2) |
| СЗВ—НЗВ | 0.959 (19) | C16—H16B | 0.965 (18) |
| C4B—C5B | 1.390 (2) | C16—H16A | 0.99 (2) |

| C4B—N4B | 1.446 (2) | | |
|--|---------------------------|---------------------|-------------------------|
| | | | |
| O1A—C1A—C2A | 124.85 (14) | C5B—C6B—C1B | 124.82 (14) |
| O1A—C1A—C6A | 123.25 (15) | C5B—C6B—N6B | 116.40 (13) |
| C2A—C1A—C6A | 111.84 (13) | C1B—C6B—N6B | 118.76 (13) |
| C3A—C2A—C1A | 124.24 (13) | O62B—N6B—O61B | 123.87 (16) |
| C3A—C2A—N2A | 116.59 (14) | O62B—N6B—C6B | 117.52 (16) |
| C1A—C2A—N2A | 119.16 (13) | O61B—N6B—C6B | 118.50 (14) |
| O21A—N2A—O22A | 122.68 (14) | C16—N11—C12 | 110.14 (13) |
| O21A—N2A—C2A | 118.32 (14) | C16—N11—C11A | 111.97 (14) |
| O22A—N2A—C2A | 118.91 (14) | C12—N11—C11A | 111.84 (14) |
| C2A—C3A—C4A | 119.12 (15) | C16—N11—H11 | 107.5 (11) |
| С2А—С3А—Н3А | 118.9 (13) | C12—N11—H11 | 109.0 (11) |
| С4А—С3А—НЗА | 121.8 (13) | C11A—N11—H11 | 106.2 (11) |
| C3A—C4A—C5A | 121.40 (14) | N11—C11A—H11C | 106.1 (16) |
| C3A—C4A—N4A | 119.07 (14) | N11—C11A—H11B | 106.3 (15) |
| C5A—C4A—N4A | 119.46 (13) | H11C—C11A—H11B | 110 (2) |
| O42A—N4A—O41A | 123.36 (14) | N11—C11A—H11A | 106.9 (17) |
| O42A—N4A—C4A | 118.51 (14) | H11C—C11A—H11A | 116 (2) |
| O41A—N4A—C4A | 118.12 (14) | H11B—C11A—H11A | 111 (2) |
| C6A - C5A - C4A | 118.49 (14) | N11—C12—C13 | 110.48 (13) |
| C6A—C5A—H5A | 119.2 (11) | N11—C12—H12B | 106.7 (11) |
| C4A—C5A—H5A | 122.3 (11) | C13—C12—H12B | 110.9 (11) |
| C_{5A} C_{6A} C_{1A} | 124 74 (14) | N11—C12—H12A | 107.2(13) |
| C5A - C6A - N6A | 116.96 (13) | C13— $C12$ — $H12A$ | 110.5(13) |
| C1A - C6A - N6A | 118 30 (13) | H12B $C12$ $H12A$ | 110.9 (16) |
| O62A - N6A - O61A | 122.69 (14) | N14-C13-C12 | 110.29 (14) |
| O62A - N6A - C6A | 118 18 (13) | N14—C13—H13B | 107.8(11) |
| O61A - N6A - C6A | 119.09 (13) | C12— $C13$ — $H13B$ | 107.0(11) 110.5(11) |
| O1B-C1B-C2B | 124 76 (13) | N14—C13—H13A | 108.6(12) |
| O1B - C1B - C6B | 12354(14) | C12— $C13$ — $H13A$ | 110.3(12) |
| C2B-C1B-C6B | 111 65 (13) | H13B—C13—H13A | 109.3 (16) |
| C_{3B} C_{2B} C_{1B} C_{1B} | 124 68 (13) | C15-N14-C13 | 111 26 (14) |
| C_{3B} C_{2B} N_{2B} | 115.96 (14) | C15—N14—H14B | 1095(13) |
| C1B-C2B-N2B | 119.34 (13) | C13 $N14$ $H14B$ | 107.5(13) |
| O21B - N2B - O22B | 122.20 (14) | C15 $N14$ $H14A$ | 107.5(13) 108.4(13) |
| O21B N2B C2B | 118 69 (14) | C13— $N14$ — $H14A$ | 108.3(12) |
| O22B N2B C2B | 118.96 (14) | H14B 14 $H14A$ | 111.8 (18) |
| $C^{2}B - C^{3}B - C^{4}B$ | 118.77 (15) | N14— $C15$ — $C16$ | 111.0(10) 110.20(13) |
| C2B = C3B = C4B C2B = C3B = H3B | 1201(11) | N14—C15—H15B | 108.1(12) |
| C4B-C3B-H3B | 120.1(11) 121.1(11) | C16—C15—H15B | 100.1(12) 109.3(12) |
| C_{3B} C_{4B} C_{5B} C_{5B} | 121.1(11) 121.27(14) | N14-C15-H15A | 109.5(12) 108.0(13) |
| C3B - C4B - N4B | 121.27(17) 119 00 (14) | C16-C15-H15A | 110.7(13) |
| C5B - C4B - M4B | 119.73 (13) | H15B-C15-H15A | 110.7(13) 110.4(16) |
| $O42B_N4B_O41B$ | 122 80 (14) | N11_C16_ C15 | 110.7(10) |
| $O_{42}D = O_{41}D = O_{41}D$ $O_{42}B = N/B = C/B$ | 118 86 (14) | N11_C16_H16P | 100.70(13) 108.1(10) |
| $O_{4}D$ O | 118.00(14) 118.24(14) | C15 $C16$ $H16B$ | 112 1 (10) |
| C_{1D} C_{1D} C_{1D} C_{1D} C_{1D} | 110.24 (14) | N11 C16 H16A | 112.1(11) 108.0(12) |
| COD-CJD-CHD | 110.00(14) | 1111-CIU-1110A | 100.7(13) |

| C6B—C5B—H5B | 120.0 (11) | C15—C16—H16A | 108.8 (13) |
|------------------|--------------|------------------|--------------|
| C4B—C5B—H5B | 121.3 (11) | H16B—C16—H16A | 108.2 (16) |
| | | | |
| O1A—C1A—C2A—C3A | 172.55 (16) | C3B—C2B—N2B—O21B | 27.4 (2) |
| C6A—C1A—C2A—C3A | -4.8 (2) | C1B—C2B—N2B—O21B | -153.91 (17) |
| O1A—C1A—C2A—N2A | -8.5 (2) | C3B—C2B—N2B—O22B | -148.21 (17) |
| C6A—C1A—C2A—N2A | 174.11 (14) | C1B—C2B—N2B—O22B | 30.5 (2) |
| C3A—C2A—N2A—O21A | -26.5 (2) | C1B—C2B—C3B—C4B | -2.8 (2) |
| C1A—C2A—N2A—O21A | 154.53 (16) | N2B-C2B-C3B-C4B | 175.80 (14) |
| C3A—C2A—N2A—O22A | 150.03 (16) | C2B—C3B—C4B—C5B | -0.3 (2) |
| C1A—C2A—N2A—O22A | -29.0 (2) | C2B—C3B—C4B—N4B | -179.82 (14) |
| C1A—C2A—C3A—C4A | 4.6 (2) | C3B—C4B—N4B—O42B | 3.6 (2) |
| N2A—C2A—C3A—C4A | -174.39 (14) | C5B—C4B—N4B—O42B | -175.90 (16) |
| C2A—C3A—C4A—C5A | -1.8 (2) | C3B—C4B—N4B—O41B | -176.40 (15) |
| C2A—C3A—C4A—N4A | 175.04 (14) | C5B—C4B—N4B—O41B | 4.1 (2) |
| C3A—C4A—N4A—O42A | 15.4 (2) | C3B—C4B—C5B—C6B | 0.3 (2) |
| C5A—C4A—N4A—O42A | -167.73 (15) | N4B—C4B—C5B—C6B | 179.78 (14) |
| C3A—C4A—N4A—O41A | -163.68 (14) | C4B—C5B—C6B—C1B | 2.9 (2) |
| C5A—C4A—N4A—O41A | 13.2 (2) | C4B-C5B-C6B-N6B | -175.30 (14) |
| C3A—C4A—C5A—C6A | -0.2 (2) | O1B-C1B-C6B-C5B | 172.05 (15) |
| N4A—C4A—C5A—C6A | -177.01 (14) | C2B—C1B—C6B—C5B | -5.4 (2) |
| C4A—C5A—C6A—C1A | -0.4 (2) | O1B-C1B-C6B-N6B | -9.8 (2) |
| C4A—C5A—C6A—N6A | 179.97 (13) | C2B-C1B-C6B-N6B | 172.77 (14) |
| O1A—C1A—C6A—C5A | -174.70 (15) | C5B—C6B—N6B—O62B | -38.6 (2) |
| C2A—C1A—C6A—C5A | 2.7 (2) | C1B—C6B—N6B—O62B | 143.11 (16) |
| O1A—C1A—C6A—N6A | 4.9 (2) | C5B—C6B—N6B—O61B | 137.68 (16) |
| C2A—C1A—C6A—N6A | -177.70 (13) | C1B—C6B—N6B—O61B | -40.6 (2) |
| C5A—C6A—N6A—O62A | 33.9 (2) | C16—N11—C12—C13 | -58.20 (17) |
| C1A—C6A—N6A—O62A | -145.77 (16) | C11A—N11—C12—C13 | 176.60 (15) |
| C5A—C6A—N6A—O61A | -144.12 (17) | N11-C12-C13-N14 | 57.33 (18) |
| C1A—C6A—N6A—O61A | 36.3 (2) | C12-C13-N14-C15 | -56.74 (18) |
| O1B—C1B—C2B—C3B | -172.05 (15) | C13—N14—C15—C16 | 56.51 (19) |
| C6B—C1B—C2B—C3B | 5.3 (2) | C12—N11—C16—C15 | 58.16 (17) |
| O1B—C1B—C2B—N2B | 9.4 (2) | C11A—N11—C16—C15 | -176.72 (14) |
| C6B—C1B—C2B—N2B | -173.25 (14) | N14—C15—C16—N11 | -57.15 (19) |
| | | | |

Hydrogen-bond geometry (Å, °)

| D—H···A | D—H | Н…А | D····A | D—H…A |
|---|------------|------------|-------------|------------|
| N11—H11…O1A | 0.892 (18) | 1.831 (18) | 2.6305 (17) | 148.1 (16) |
| N11—H11···O22A | 0.892 (18) | 2.356 (18) | 2.996 (2) | 128.8 (14) |
| N14—H14 B ···O1 B^{i} | 0.88 (2) | 1.98 (2) | 2.8181 (19) | 157.3 (17) |
| N14—H14 <i>A</i> ···O1 <i>B</i> | 0.92 (2) | 1.99 (2) | 2.7962 (18) | 146.4 (18) |
| N14—H14 <i>A</i> ···O22 <i>B</i> | 0.92 (2) | 2.28 (2) | 2.992 (2) | 133.9 (16) |
| C5 <i>A</i> —H5 <i>A</i> ···O21 <i>A</i> ⁱⁱ | 0.917 (19) | 2.476 (19) | 3.383 (2) | 170.3 (16) |
| C5 <i>B</i> —H5 <i>B</i> ···O21 <i>B</i> ⁱⁱⁱ | 0.913 (18) | 2.487 (18) | 3.394 (2) | 172.3 (15) |
| C11 <i>A</i> —H11 <i>C</i> ···O41 <i>A</i> ^{iv} | 0.93 (3) | 2.48 (3) | 3.345 (2) | 155 (2) |
| C11 <i>A</i> —H11 <i>A</i> ···O62 <i>A</i> ⁱⁱⁱ | 0.94 (3) | 2.57 (3) | 3.496 (3) | 168 (2) |
| | | | | |

supporting information

| C12—H12A····O22A | 0.96 (2) | 2.56 (2) | 3.046 (2) | 111.6 (15) |
|----------------------------|----------|----------|-----------|------------|
| C13—H13 A ···O62 B^{v} | 0.96 (2) | 2.46 (2) | 3.386 (2) | 162.9 (17) |

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