

Acta Crystallographica Section C Structural Chemistry ISSN 2053-2296

Three-dimensional hydrogen-bonded framework structures in flunarizinium nicotinate and flunarizinediium bis(4-toluenesulfonate) dihydrate

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Received 15 July 2014 Accepted 16 July 2014

The structures of two salts of flunarizine, namely 1-bis[(4fluorophenyl)methyl]-4-[(2E)-3-phenylprop-2-en-1-yl]piperazine, C₂₆H₂₆F₂N₂, are reported. In flunarizinium nicotinate {systematic name: 4-bis[(4-fluorophenyl)methyl]-1-[(2E)-3phenylprop-2-en-1-yl]piperazin-1-ium pyridine-3-carboxylate}, $C_{26}H_{27}F_2N_2^+ \cdot C_6H_4NO_2^-$, (I), the two ionic components are linked by a short charge-assisted N-H···O hydrogen bond. The ion pairs are linked into a three-dimensional framework structure by three independent $C-H \cdots O$ hydrogen bonds, augmented by $C-H \cdot \cdot \pi$ (arene) hydrogen bonds and an aromatic π - π stacking interaction. In flunarizinediium bis(4-toluenesulfonate) dihydrate {systematic name: 1-[bis(4-fluorophenyl)methyl]-4-[(2E)-3-phenylprop-2en-1-yl]piperazine-1,4-diium bis(4-methylbenzenesulfonate) dihydrate}, C₂₆H₂₈F₂N₂²⁺·2C₇H₇O₃S⁻·2H₂O, (II), one of the anions is disordered over two sites with occupancies of 0.832 (6) and 0.168 (6). The five independent components are linked into ribbons by two independent N-H···O hydrogen bonds and four independent O-H···O hydrogen bonds, and these ribbons are linked to form a three-dimensional framework by two independent $C-H \cdots O$ hydrogen bonds, but C-H··· π (arene) hydrogen bonds and aromatic π - π stacking interactions are absent from the structure of (II). Comparisons are made with some related structures.

Keywords: crystal structure; flunarizinediium salt; flunarizinium salt; hydrogen bonding; π - π stacking; calcium-channel blocker.

1. Introduction

Flunarizine {systematic name: $1-[bis(4-fluorophenyl)methyl]-4-[(2E)-3-phenylprop-2-en-1-yl]piperazine} is a nonselective$



calcium-channel blocker (Amery, 1983; Fagbemi et al., 1984; Tarland & Flatmark, 1999) which is effective in the prophylaxis of migraine, occlusive peripheral vascular disease and vertigo of central and peripheral origin, and also as an adjuvant in the therapy of epilepsy. Its pharmacodynamic and pharmacokinetic properties and its therapeutic use have been reviewed (Holmes et al., 1984). Brief reports on the structures of several salts derived from flunarizine (Kavitha, Jasinski et al., 2013; Kavitha, Yathirajan et al., 2013; Shivaprakash et al., 2014) and from substituted piperazine derivatives closely related to flunarizine (Kavitha, Butcher et al., 2013; Kavitha, Yildirim et al., 2013) have been published recently, and we report here the molecular and supramolecular structures of flunarizinium nicotinate, (I) (Fig. 1), and flunarizinediium bis(4-toluenesulfonate) dihydrate, (II) (Fig. 2). The main purposes of the present study are, firstly, to compare the supramolecular assemblies of (I) and (II), and, secondly, to compare the structures of (I) and (II) with those of some closely related analogues.



2. Experimental

2.1. Synthesis and crystallization

For the synthesis of salt (I), flunarizine free base (4.05 g, 0.01 mol) and nicotinic acid (1.23 g, 0.01 mol) were dissolved in hot dimethylformamide (5 ml) and the solution was stirred for 10 min. The solution was then allowed to cool slowly to ambient temperature in the presence of air. Colourless crystals of (I) suitable for single-crystal X-ray diffraction appeared after a few days and were collected by filtration (m.p. 383–

Table 1

Experimental details.

| | (I) | (II) |
|--|---|---|
| Crystal data | | |
| Chemical formula | $C_{26}H_{27}F_2N_2^+ \cdot C_6H_4NO_2^-$ | $C_{26}H_{28}F_2N_2^{2+}\cdot 2C_7H_7O_3S^{-}\cdot 2H_2O$ |
| $M_{\rm r}$ | 527.60 | 784.91 |
| Crystal system, space group | Monoclinic, Pc | Monoclinic, $P2_1/c$ |
| Temperature (K) | 200 | 200 |
| a, b, c (Å) | 10.8536 (4), 10.8103 (4), 11.3901 (4) | 10.0546 (5), 14.8338 (6), 26.9437 (12) |
| β (°) | 92.717 (2) | 106.497 (3) |
| $V(Å^3)$ | 1334.91 (8) | 3853.2 (3) |
| Ζ | 2 | 4 |
| Radiation type | Μο Κα | Μο Κα |
| $\mu \text{ (mm}^{-1})$ | 0.09 | 0.20 |
| Crystal size (mm) | $0.54 \times 0.49 \times 0.13$ | $0.59 \times 0.43 \times 0.24$ |
| Data collection | | |
| Diffractometer | Bruker APEXII CCD area-detector diffractometer | Bruker APEXII CCD area-detector diffractometer |
| Absorption correction | Multi-scan (SADABS; Sheldrick, 2003) | Multi-scan (SADABS; Sheldrick, 2003) |
| T_{\min}, T_{\max} | 0.873, 0.988 | 0.877, 0.953 |
| No. of measured, independent and observed $[I > 2\sigma(I)]$ reflections | 23278, 6092, 5770 | 45646, 7212, 6219 |
| R _{int} | 0.027 | 0.038 |
| $(\sin \theta / \lambda)_{\rm max} ({\rm \AA}^{-1})$ | 0.668 | 0.607 |
| Refinement | | |
| $R[F^2 > 2\sigma(F^2)], wR(F^2), S$ | 0.030, 0.081, 1.03 | 0.075, 0.202, 1.09 |
| No. of reflections | 6092 | 7212 |
| No. of parameters | 352 | 527 |
| No. of restraints | 2 | 27 |
| H-atom treatment | H-atom parameters constrained | H-atom parameters constrained |
| $\Delta \rho_{\rm max}, \Delta \rho_{\rm min} \ ({\rm e} \ {\rm \AA}^{-3})$ | 0.15, -0.18 | 0.71, -0.60 |
| Absolute structure | Flack x parameter (Flack, 1983) determined using 2539 quotients $[(I^+) - (I^-)]/[(I^+) + (I^-)]$ (Parsons et al. 2013) | - |
| Absolute structure parameter | 0.10 (15) | _ |
| 1 | | |

Computer programs: APEX2 (Bruker, 2010), SAINT (Bruker, 2010), SHELXS97 (Sheldrick, 2008), SHELXL2014 (Sheldrick, 2014) and PLATON (Spek, 2009).

385 K). For the synthesis of hydrated salt (II), flunarizine free base (4.05 g, 0.01 mol) and 4-toluenesufonic acid mono-hydrate (1.72 g, 0.009 mol) were dissolved in hot methanol



Figure 1

The independent components of salt (I), showing the atom-labelling scheme and the $N-H\cdots O$ hydrogen bond (dashed line) within the selected asymmetric unit. Displacement ellipsoids are drawn at the 30% probability level.

(20 ml) and the solution was stirred for 10 min. The solution was then allowed to cool slowly to ambient temperature in the presence of air. Yellow crystals of (II) suitable for single-crystal X-ray diffraction appeared after a few days and were collected by filtration (m.p. 406–410 K).

2.2. Refinement

Crystal data, data collection and structure refinement details are summarized in Table 1. All H atoms were located in difference maps and then treated as riding atoms. C-bound H atoms were treated as riding in geometrically idealized positions, with C-H = 0.95 (alkenyl and aromatic), 0.98 (CH₃), 0.99 (CH₂) or 1.00 Å (aliphatic CH), and with $U_{iso}(H) =$ $kU_{eq}(C)$, where k = 1.5 for the methyl groups, which were permitted to rotate but not to tilt, and 1.2 for all other C-bound H atoms. N- and O-bound H atoms were permitted to ride at the positions located in difference maps, with $U_{iso}(H) = 1.2U_{eq}(N)$ or $1.5U_{eq}(O)$, giving the N-H and O-H distances shown in Tables 2 and 3. For (I), the correct orientation of the structure with respect to the polar-axis direction was established by means of the Flack x parameter (Flack, 1983), determined as x = 0.09 (15) by the use of 2539 quotients $[(I^+) - (I^-)]/[(I^+) + (I^-)]$ (Parsons *et al.*, 2013) from a total of 2758 Bijvoet pairs (83% coverage), and by the use of the Hooft y parameter (Hooft et al., 2008), calculated as 0.10 (17), also using 2758 Bijvoet pairs. For hydrated salt (II), the crys-



Figure 2

The independent components of hydrated salt (II), showing the atom-labelling scheme and the $O-H \cdots O$ and $N-H \cdots O$ hydrogen bonds (dashed lines) within the selected asymmetric unit. Displacement ellipsoids are drawn at the 30% probability level and the site occupancies for the two orientations of the disordered anion are 0.832 (6) and 0.168 (6). The pairs of atomic sites S61 and S61A, and O63 and O63A, are nearly coincident and, for the sake of clarity, the atom labels C61A–C67A, S61A and O63A have been omitted.

tals were all of very indifferent quality and about a dozen individual crystals were subjected to preliminary examination using polarizing microscopy; none showed any visible signs of twinning or intergrowth. Three data sets were obtained from the two most promising looking crystals and the refinement reported here results from the best of these data sets, although all three gave essentially identical solutions. For all three data sets, it was apparent from an early stage that one of the anions, that containing atom S61 (Fig. 2), was disordered over two sets of sites. For the minor orientation, the directly bonded distances and the one-angle distances were restrained to be equal to the corresponding distances in the major orientation, subject to standard uncertainties of 0.005 and 0.01 Å, respectively. In addition, the anisotropic displacement parameters for the corresponding pairs of partial-occupancy atoms which occupied approximately the same regions of physical

Table 2

Hydrogen-bond geometry (Å, °) for (I).

Cg1 and Cg2 represent the centroids of the C11–C16 and C31–C36 rings, respectively.

| $D - H \cdots A$ | D-H | $H \cdot \cdot \cdot A$ | $D \cdot \cdot \cdot A$ | $D - \mathbf{H} \cdot \cdot \cdot A$ |
|------------------------------|------|-------------------------|-------------------------|--------------------------------------|
| N4-H4···O51 | 0.98 | 1.58 | 2.5630 (19) | 175 |
| $C3-H3B\cdots O52^{i}$ | 0.99 | 2.53 | 3.440 (2) | 153 |
| C13-H13···O52 ⁱⁱ | 0.95 | 2.57 | 3.442 (2) | 152 |
| C56-H56···O51 ⁱⁱⁱ | 0.95 | 2.34 | 3.236 (3) | 158 |
| $C26-H26\cdots Cg1^{iv}$ | 0.95 | 2.81 | 3.758 (2) | 173 |
| $C55-H55\cdots Cg2^{v}$ | 0.95 | 2.85 | 3.567 (2) | 133 |

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

space were constrained to be identical. Subject to these conditions, the site occupancies for the two orientations refined to 0.832 (6) and 0.168 (6). Because of the rather weak diffraction at high angles, all reflections having $\theta > 25.5^{\circ}$ were omitted from the final refinements for hydrated salt (II). Examination of the refined structures of (I) and (II) using *PLATON* (Spek, 2009) showed an absence of any solvent-accessible voids for both structures. For the rerefinement of (VI) (see *Results and discussion*, §3), the reported unit cell in the space group $P2_1/n$ (Kavitha, Butcher *et al.*, 2013) was transformed to the $P2_1/c$ setting with dimensions a = 10.0845 (2) Å, b = 14.6026 (3) Å, c = 27.1907 (7) Å and $\beta = 108.315$ (2)°.

Table 3Hydrogen-bond geometry (Å, $^{\circ}$) for (II)

| $D - H \cdots A$ | D-H | $H \cdot \cdot \cdot A$ | $D \cdots A$ | $D - H \cdots A$ | |
|---|------|-------------------------|--------------|------------------|--|
| N1-H1071 | 0.93 | 1 79 | 2 716 (4) | 169 | |
| N4-H4···O81 | 1.00 | 1.69 | 2.690(4) | 173 | |
| O71−H71A···O51 | 0.85 | 1.88 | 2.708 (4) | 167 | |
| $O71 - H71B \cdot \cdot \cdot O61^{i}$ | 0.84 | 1.87 | 2.675 (5) | 160 | |
| O81−H81A···O61 | 0.84 | 2.05 | 2.852 (5) | 161 | |
| $O81 - H81B \cdot \cdot \cdot O52^{ii}$ | 0.84 | 1.92 | 2.748 (4) | 167 | |
| $C1-H1A\cdots O63$ | 1.00 | 2.23 | 3.139 (7) | 151 | |
| $C2-H2B\cdots O63$ | 0.99 | 2.47 | 3.314 (7) | 143 | |
| $C6-H6A\cdots O53^{iii}$ | 0.99 | 2.37 | 3.191 (5) | 140 | |
| $C16-H16\cdots O62^{i}$ | 0.95 | 2.41 | 3.228 (6) | 144 | |
| C22-H22···O71 | 0.95 | 2.49 | 3.381 (5) | 156 | |
| C34-H34···O51 ^{iv} | 0.95 | 2.42 | 3.362 (6) | 169 | |
| C41-H41A···O53 | 0.99 | 2.35 | 3.280 (5) | 157 | |
| $C41 - H41B \cdots O61^{v}$ | 0.99 | 2.37 | 3.341 (5) | 167 | |

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

3. Results and discussion

Salt (I) consists of flunarizinium monocations, in which protonation of the flunarizine free base has occurred selectively at atom N4, which is the N atom more remote from the electronegative fluorophenyl units, and nicotinate anions; the asymmetric unit was selected such that the two ions within it are linked by an N-H···O hydrogen bond (Fig. 1 and Table 2). The constitution of (II) is more complex, comprising a flunarizinediium dication, two 4-methylbenzenesulfonate anions, one of which exhibits positional disorder, and two water molecules. Accordingly, there is considerable flexibility in the choice of the asymmetric unit. However, it is possible to specify a reasonably compact asymmetric unit within which the five independent components are linked by N-H···O and O-H···O hydrogen bonds (Fig. 2 and Table 3).

In each of (I) and (II), the pyrazine ring adopts a chair conformation, with the two hydrocarbyl substituents occupying equatorial sites (Figs. 1 and 2). The ring-puckering parameters (Cremer & Pople, 1975), calculated for the atom sequence N1–C2–C3–N4–C5–C6, are Q = 0.5934 (17) Å, $\theta = 3.56$ (15)° and $\varphi = 345$ (3)° for (I), and Q = 0.554 (3) Å, $\theta = 3.0$ (3)° and $\varphi = 326$ (8)° for (II). For an ideal chair conformation, the value of θ is exactly zero (Boeyens, 1978).

The 3-phenylprop-2-en-1-yl substituents in (I) and (II) adopt very similar orientations relative to the piperazine ring and very similar overall conformations, as shown by the relevant torsion angles (Tables 4 and 5). On the other hand, the orientations of the two independent fluorophenyl rings are somewhat different in the two compounds. This may be ascribed, at least in part, to the different direction-specific interactions involving these rings in the two compounds, including the C-H···O and C-H··· π (arene) hydrogen bonds and the aromatic π - π stacking interactions, which differ markedly between the two compounds, as discussed below. There is no internal symmetry in the cation in either compound, so that these cations are conformationally chiral. However, in each case, the presence of glide planes confirms that equal numbers of the two conformational enantiomers are present.

Within the selected asymmetric unit of (I) (Fig. 1), the two components are linked by a rather short and nearly linear charge-assisted (Gilli *et al.*, 1994) $N-H\cdots O$ hydrogen bond

| Ta | ble | 4 |
|----|-----|---|
|----|-----|---|

| Selected torsion | angles (| (°) |) for | (I) | 6 |
|------------------|----------|-----|-------|-----|---|
|------------------|----------|-----|-------|-----|---|

| C2-N1-C1-C11 | -70.92 (16) | C3-N4-C41-C42 | -55.58 (19) |
|---------------|-------------|-----------------|--------------|
| C2-N1-C1-C21 | 165.22 (13) | N4-C41-C42-C43 | 118.27 (19) |
| N1-C1-C11-C12 | 150.17 (15) | C41-C42-C43-C31 | 177.23 (16) |
| N1-C1-C21-C22 | 69.4 (2) | C42-C43-C31-C32 | -178.93 (18) |
| | | | |

| Table 5 | | | | | |
|----------|---------|--------|-----|-----|-------|
| Selected | torsion | angles | (°) | for | (II). |

| | ., ., | | |
|---------------|-----------|-----------------|------------|
| C2-N1-C1-C11 | -51.9 (4) | C3-N4-C41-C42 | -56.2(4) |
| C2-N1-C1-C21 | -179.0(3) | N4-C41-C42-C43 | 121.6 (4) |
| N1-C1-C11-C12 | 101.8 (4) | C41-C42-C43-C31 | 172.9 (3) |
| N1-C1-C21-C22 | 42.4 (5) | C42-C43-C31-C32 | -167.1 (4) |



Figure 3

Part of the crystal structure of salt (I), showing the formation of a hydrogen-bonded C(7) chain parallel to [001] and built from nicotinate anions only. Hydrogen bonds are shown as dashed lines and, for the sake of clarity, H atoms not involved in the motif shown have been omitted. Atoms marked with an asterisk or a hash symbol (#) are at the symmetry positions $(x, -y + 2, z - \frac{1}{2})$ and $(x, -y + 2, z + \frac{1}{2})$, respectively.

(Table 2). Ion pairs of this type are linked by three independent $C-H \cdots O$ hydrogen bonds (Table 2) to form a threedimensional framework structure, the formation of which is



Figure 4

A stereoview of part of the crystal structure of salt (I), showing the formation of a hydrogen-bonded $C_2^2(7)$ chain parallel to [001] and containing alternating C-H···O and N-H···O hydrogen bonds, shown as dashed lines. For the sake of clarity, H atoms bonded to C atoms which are not involved in the motif shown have been omitted.





Figure 5

A stereoview of part of the crystal structure of salt (I), showing the formation of a hydrogen-bonded $C_2^2(13)$ chain parallel to $[20\overline{1}]$ and containing alternating $C-H\cdots O$ and $N-H\cdots O$ hydrogen bonds, shown as dashed lines. For the sake of clarity, H atoms bonded to C atoms which are not involved in the motif shown have been omitted.

readily analysed in terms of three fairly simple one-dimensional substructures (Ferguson et al., 1998a,b; Gregson et al., 2000). The simplest of these substructures involves only the nicotinate component, where anions related by the *c*-glide plane at y = 1 are linked into a C(7) chain (Bernstein *et al.*, 1995) running parallel to the [001] direction (Fig. 3). The C- $H \cdots O$ hydrogen bond having atom C3 as the donor, together with the $N-H \cdots O$ hydrogen bond, link ion pairs related by the *c*-glide plane at $y = \frac{1}{2}$ to form a $C_2^2(7)$ chain, also running parallel to the [001] direction (Fig. 4). The combination of the two chain motifs parallel to [001] generates a sheet lying parallel to (100) and such sheets are linked by the third onedimensional substructure. In this final substructure, the C- $H \cdots O$ hydrogen bond having atom C13 as the donor links ion pairs, again related by the *c*-glide plane at $y = \frac{1}{2}$, but this time forming a $C_2^2(13)$ chain running parallel to the [201] direction (Fig. 5), so completing the formation of the three-dimensional structure.

The resulting framework of (I) is modestly reinforced by two fairly long C-H··· π (arene) hydrogen bonds (Table 2) and by an aromatic π - π stacking interaction. Between the C21-C26 ring in the cation at (x, y, z) and the C31-C36 ring in the cation at $(-1 + x, 1 - y, -\frac{1}{2} + z)$ there is a ring-centroid separation of 3.6986 (11) Å. The dihedral angle between the ring planes is 5.02 (9)° and the shortest perpendicular distance from the centroid of one ring to the plane of the other is 3.3875 (8) Å, corresponding to a nearly ideal ring-centroid offset of *ca* 1.48 Å.

In (II), the five independent components are again linked into a three-dimensional framework structure by a combination of $O-H\cdots O$, $N-H\cdots O$ and $C-H\cdots O$ hydrogen bonds





A stereoview of part of the crystal structure of hydrated salt (II), showing the formation of a hydrogen-bonded ribbon parallel to [100]. Hydrogen bonds are shown as dashed lines and, for the sake of clarity, H atoms not involved in the motif shown have been omitted and only the major orientation of the disordered anion is shown.

(Table 3), although $C-H\cdots\pi(\text{arene})$ hydrogen bonds and aromatic $\pi-\pi$ stacking interactions are absent from the structure. The framework of (II) is considerably more complex than that of (I) but, as for (I), its formation can readily be analysed in terms of low-dimensional substructures. The principal substructure involves only $O-H\cdotsO$ and N- $H\cdotsO$ hydrogen bonds. The five-component aggregates (Fig. 2), which are related by translation along [100], are linked to form a broad ribbon in which the cations, acting as twofold donors in $N-H\cdotsO$ hydrogen bonds, alternate with $R_4^3(10)$ rings built from the two independent anions and the two independent water molecules (Fig. 6). Four of these ribbons pass through each unit cell, with the hydrogen bonds lying approximately along the lines (x, 0.4, 0.1), (x, 0.9, 0.4), (x,0.6, 0.9) and (x, 0.1, 0.6).

The structure of (II) contains a number of short intermolecular $C-H\cdots O$ contacts (Table 3). However, the majority of these lie either within the selected asymmetric unit or within the ribbon built from the $O-H\cdots O$ and $N-H\cdots O$ hydrogen bonds, or they have $C-H\cdots O$ angles close to 140°, such that they cannot be regarded as structurally significant (Wood *et al.*, 2009). Hence, the only two of these interactions which can be regarded as structurally significant hydrogen bonds influencing the overall dimensionality of the supramolecular assembly are those having atoms O51 and O61 as the acceptors, and each of these hydrogen bonds can be regarded as the basis of a simple substructure.

The C-H···O hydrogen bond involving atom O51 leads to the formation of a $C_3^2(16)$ chain running parallel to the [010] direction, comprising only the cation, the anion containing atom S51 and the water molecule containing atom O71, and built from components related by the 2_1 screw axis along $(0, y, \frac{1}{4})$ (Fig. 7). The effect of this chain motif is to link the [100] chains in the domain $0 < z < \frac{1}{2}$ into a sheet lying parallel to (001). A second sheet of this type, related to the first by inversion, lies in the domain $\frac{1}{2} < z < 1.0$. The final substructure involves, in addition to the cation, the anion containing atom S61 and the water molecule containing atom O81 (*i.e.* those



Figure 7

A stereoview of part of the crystal structure of hydrated salt (II), showing the formation of a hydrogen-bonded $C_3^2(16)$ chain parallel to [010] and incorporating the cation and only one type each of the independent anions and water molecule. Hydrogen bonds are shown as dashed lines and, for the sake of clarity, H atoms not involved in the motif shown have been omitted.

not participating in the chain along [010]), and it takes the form of a finite zero-dimensional motif characterized by an $R_6^4(14)$ ring (Fig. 8). The effect of this ring motif is to link directly the sheet in the domain $0 < z < \frac{1}{2}$ with two adjacent sheets in the domains $\frac{1}{2} < z < 1.0$ and $-\frac{1}{2} < z < 0$, so completing the formation of the three-dimensional framework structure.

It is of interest briefly to compare the supramolecular assembly in some closely-related compounds with that reported here for (I) and (II). Flunarizine forms 1:1 salts with both succinic and maleic acids, and in flunarizinium hydrogen succinate, (III) (Kavitha, Yathirajan et al., 2013), the anion adopts an extended chain conformation such that the carboxylic acid -OH group is available as a donor in the formation of inter-anion O-H···O hydrogen bonds. By contrast, the anion in flunarizinium hydrogen maleate, (IV) (Kavitha, Jasinski et al., 2013), contains a short intra-anion $O-H \cdots O$ hydrogen bond, forming an S(7) motif, and the carboxylic acid -OH group is not available as a donor for the formation of hydrogen bonds with other entities in the structure. In (III), a combination of O-H···O, N-H···O and $C-H \cdots O$ hydrogen bonds links the component ions into a three-dimensional framework structure. The supramolecular structure of (IV) was described in the original report as a chain, but re-examination of this structure using the published atomic coordinates shows the structure to consist of a ribbon built from alternating edge-fused S(7) and $R_3^3(11)$ rings (Fig. 9). In flunarizinediium dichloride hemihydrate, (V) (Shivaprakash et al., 2014), the independent ionic components are linked by two nearly-linear charge-assisted N-H···Cl hydrogen bonds and symmetry-related pairs of these ion triplets are linked by $O-H \cdots Cl$ hydrogen bonds, where the donor is a water molecule lying across a twofold rotation axis,



Figure 8

Part of the crystal structure of hydrated salt (II), showing the formation of a centrosymmetric hydrogen-bonded $R_6^4(14)$ motif which links adjacent (001) sheets. Hydrogen bonds are shown as dashed lines and, for the sake of clarity, components not involved in the motif shown and H atoms bonded to C atoms not involved in the motif shown have been omitted. Atoms marked with an asterisk are at the symmetry position (-x + 1, -y + 1, -z).

to form a seven-component aggregate. Two independent and nearly linear C—H···Cl hydrogen bonds link these aggregates into a complex ribbon. There are also two short C—H···O contacts in the structure of (V), but in each the H···O distance (2.67 and 2.68 Å) exceeds the sum of the van der Waals radii (2.61 Å; Bondi, 1964; Rowland & Taylor, 1996) while having C—H···O angles close to 140° (cf. Wood et al., 2009), so that neither of these contacts is structurally significant.



Figure 9

A stereoview of part of the crystal structure of (IV), showing the formation of a hydrogen-bonded ribbon containing alternating S(7) and $R_3^3(11)$ rings. The original atomic coordinates (Kavitha, Jasinski *et al.*, 2013) have been used. Hydrogen bonds are shown as dashed lines and, for the sake of clarity, H atoms not involved in the motifs shown have been omitted.

Cinnarizine {systematic name: 1-diphenylmethyl-4-[(2E)-3phenylprop-2-en-1-ylpiperazine} is also a calcium-channel blocker, which differs from flunarizine only in the absence of the 4-fluoro substituents in the diarylmethyl unit. Cinnarizine forms a dihydrated 1:2 salt, (VI), with 4-toluenesulfonic acid (Kavitha, Butcher et al., 2013). Compound (VI) appears to be isomorphous with (II), although it was refined in the alternative $P2_1/n$ setting, rather than in $P2_1/c$ as for (II). There is very little discussion of the supramolecular assembly in the original report, apart from a listing of the $O-H \cdots O$ and N- $H \cdots O$ hydrogen bonds, while $C - H \cdots O$ hydrogen bonds were not mentioned. Re-examination of the supramolecular assembly of (VI) using the published atomic coordinates shows that it contains the same type of ribbon along [100], built from O-H···O and N-H···O hydrogen bonds, as found here for (II) (cf. Fig. 6). In addition, there are two significant $C-H \cdots O$ hydrogen bonds in the structure of (VI), one of which generates a $C_3^2(16)$ chain running parallel to [010], while the other generates a centrosymmetric $R_6^4(14)$ motif, entirely comparable with the action of the corresponding hydrogen bonds in (II) (cf. Figs. 7 and 8). Hence (II) and (VI) are isostructural, as confirmed by a new refinement for (VI) carried out in the space group $P2_1/c$ using the deposited structure-factor data. This refinement, using the coordinates of (II) as the starting point, but with the two F atoms replaced by H atoms, converged to R = 0.0197 for 7891 observed reflections and 509 parameters subject to 27 restraints, with refined site occupancies for the disordered anion of 0.804 (2) and 0.196 (2), thus demonstrating clearly the isostructural nature of (II) and (VI). It is noteworthy that the absence of F substituents from (VI) appears to have no significant influence on the structure, apart from the minor and expected difference in the unit-cell volumes for (II) and (VI). *viz.* 3853.2 (3) $Å^3$ at 200 (2) K and 3801.25 (14) $Å^3$ at 100 (2) K, respectively.

CNK thanks the University of Mysore for research facilities and is grateful to the Principal, Maharani's Science College for Women, Mysore, for giving permission to carry out this research.

Supporting information for this paper is available from the IUCr electronic archives (Reference: SK3553).

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supporting information

Acta Cryst. (2014). C70, 805-811 [doi:10.1107/S2053229614016532]

Three-dimensional hydrogen-bonded framework structures in flunarizinium nicotinate and flunarizinediium bis(4-toluenesulfonate) dihydrate

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Computing details

For both compounds, data collection: *APEX2* (Bruker, 2010); cell refinement: *SAINT* (Bruker, 2010); data reduction: *SAINT* (Bruker, 2010); program(s) used to solve structure: *SHELXS97* (Sheldrick, 2008); program(s) used to refine structure: *SHELXL2014* (Sheldrick, 2014); molecular graphics: *PLATON* (Spek, 2009); software used to prepare material for publication: *SHELXL2014* (Sheldrick, 2014) and *PLATON* (Spek, 2009).

(I) 4-Bis[(4-fluorophenyl)methyl]-1-[(2E)-3-phenylprop-2-en-1-yl]piperazin-1-ium pyridine-3-carboxylate

| Crystal data | |
|---|---|
| $C_{26}H_{27}F_2N_2\cdot C_6H_4NO_2$ $M_r = 527.60$ Monoclinic, <i>Pc</i> a = 10.8536 (4) Å b = 10.8103 (4) Å c = 11.3901 (4) Å $\beta = 92.717$ (2)° V = 1334.91 (8) Å ³ Z = 2 | F(000) = 556 $D_x = 1.313 \text{ Mg m}^{-3}$ Mo K\alpha radiation, \lambda = 0.71073 \mathbf{A} Cell parameters from 6092 reflections $\theta = 1.9-28.4^{\circ}$ $\mu = 0.09 \text{ mm}^{-1}$ T = 200 K Plate, colourless $0.54 \times 0.49 \times 0.13 \text{ mm}$ |
| Data collection Bruker APEXII CCD area-detector diffractometer φ and ω scans Absorption correction: multi-scan (<i>SADABS</i> ; Sheldrick, 2003) $T_{\min} = 0.873, T_{\max} = 0.988$ 23278 measured reflections | 6092 independent reflections 5770 reflections with $I > 2\sigma(I)$ $R_{int} = 0.027$ $\theta_{max} = 28.4^\circ, \ \theta_{min} = 1.9^\circ$ $h = -14 \rightarrow 14$ $k = -14 \rightarrow 14$ $l = -15 \rightarrow 14$ |
| RefinementRefinement on F^2 Least-squares matrix: full $R[F^2 > 2\sigma(F^2)] = 0.030$ $wR(F^2) = 0.081$ $S = 1.03$ 6092 reflections352 parameters2 restraints | Hydrogen site location: mixed H-atom parameters constrained $w = 1/[\sigma^2(F_o^2) + (0.045P)^2 + 0.1632P]$ where $P = (F_o^2 + 2F_c^2)/3$ $(\Delta/\sigma)_{max} < 0.001$ $\Delta\rho_{max} = 0.15$ e Å ⁻³ $\Delta\rho_{min} = -0.18$ e Å ⁻³ |

Absolute structure: Flack *x* parameter (Flack, 1983) determined using 2539 quotients [(I+)-(I-)]/[(I+)+(I-)] (Parsons *et al.*, 2013) Absolute structure parameter: 0.10 (15)

Special details

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

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

| | x | У | Ζ | $U_{ m iso}$ */ $U_{ m eq}$ |
|-----|---------------|--------------|---------------|-----------------------------|
| N1 | 0.17850 (12) | 0.43015 (13) | 0.31935 (12) | 0.0250 (3) |
| C2 | 0.19249 (15) | 0.55279 (15) | 0.37393 (15) | 0.0261 (3) |
| H2A | 0.1416 | 0.5572 | 0.4437 | 0.031* |
| H2B | 0.1625 | 0.6169 | 0.3175 | 0.031* |
| C3 | 0.32579 (15) | 0.57830 (16) | 0.41056 (14) | 0.0268 (3) |
| H3A | 0.3330 | 0.6617 | 0.4462 | 0.032* |
| H3B | 0.3547 | 0.5170 | 0.4703 | 0.032* |
| N4 | 0.40422 (12) | 0.57106 (13) | 0.30680 (12) | 0.0244 (3) |
| H4 | 0.3770 | 0.6362 | 0.2515 | 0.029* |
| C5 | 0.38561 (15) | 0.44918 (16) | 0.24771 (15) | 0.0276 (3) |
| H5A | 0.4157 | 0.3824 | 0.3012 | 0.033* |
| H5B | 0.4341 | 0.4463 | 0.1763 | 0.033* |
| C6 | 0.25047 (15) | 0.42811 (16) | 0.21400 (15) | 0.0273 (3) |
| H6A | 0.2206 | 0.4936 | 0.1589 | 0.033* |
| H6B | 0.2400 | 0.3472 | 0.1740 | 0.033* |
| C1 | 0.04617 (14) | 0.40285 (15) | 0.29532 (15) | 0.0250 (3) |
| H1 | 0.0071 | 0.4774 | 0.2573 | 0.030* |
| C11 | -0.01644 (14) | 0.37958 (15) | 0.41020 (15) | 0.0251 (3) |
| C12 | -0.14102 (15) | 0.40862 (16) | 0.41860 (16) | 0.0302 (4) |
| H12 | -0.1844 | 0.4472 | 0.3541 | 0.036* |
| C13 | -0.20273 (17) | 0.38200 (18) | 0.51973 (18) | 0.0358 (4) |
| H13 | -0.2879 | 0.4003 | 0.5247 | 0.043* |
| C14 | -0.13720 (18) | 0.32879 (16) | 0.61156 (17) | 0.0345 (4) |
| F14 | -0.19601 (13) | 0.30702 (13) | 0.71334 (11) | 0.0516 (3) |
| C15 | -0.01478 (18) | 0.29758 (17) | 0.60796 (17) | 0.0336 (4) |
| H15 | 0.0275 | 0.2594 | 0.6733 | 0.040* |
| C16 | 0.04540 (16) | 0.32364 (16) | 0.50559 (16) | 0.0294 (3) |
| H16 | 0.1301 | 0.3028 | 0.5009 | 0.035* |
| C21 | 0.02675 (15) | 0.29468 (16) | 0.21188 (14) | 0.0265 (3) |
| C22 | 0.05734 (17) | 0.17386 (17) | 0.24553 (18) | 0.0346 (4) |
| H22 | 0.0922 | 0.1585 | 0.3222 | 0.041* |
| C23 | 0.0372 (2) | 0.07615 (19) | 0.1681 (2) | 0.0435 (5) |
| H23 | 0.0577 | -0.0061 | 0.1911 | 0.052* |
| C24 | -0.01274 (19) | 0.1004 (2) | 0.0577 (2) | 0.0454 (5) |
| F24 | -0.03385 (15) | 0.00390 (16) | -0.01654 (15) | 0.0702 (5) |

| H25 -0.0782 0.2311 -0.0565 0.052^{*} C26 $-0.02378 (17)$ $0.31482 (19)$ $0.09868 (16)$ $0.0334 (4)$ H26 -0.0449 0.3965 0.0745 0.040^{*} C41 $0.53859 (16)$ $0.58793 (18)$ $0.34023 (17)$ $0.0321 (4)$ H41A 0.5671 0.5174 0.3899 0.039^{*} C42 $0.56485 (16)$ $0.70599 (18)$ $0.40543 (17)$ $0.0327 (4)$ H42 0.5461 0.7823 0.3673 0.039^{*} C43 $0.61233 (16)$ $0.70778 (17)$ $0.51436 (16)$ $0.0313 (4)$ H43 0.6269 0.6293 0.5502 0.038^{*} C31 $0.64836 (15)$ $0.81561 (17)$ $0.58665 (15)$ $0.0299 (4)$ C32 $0.70012 (17)$ $0.77671 (19)$ $0.69949 (17)$ $0.0342 (4)$ H32 0.7088 0.7147 0.7287 0.041^{*} C33 $0.73911 (18)$ $0.8497 (2)$ $0.77006 (18)$ $0.0411 (4)$ H33 0.7747 0.8797 0.8466 0.048^{*} C34 $0.72614 (19)$ $1.0147 (2)$ $0.7288 (2)$ $0.0425 (4)$ H34 0.7532 1.0824 $0.7760 (18)$ $0.0417 (4)$ H35 0.6637 1.1180 0.5893 0.050^{*} C36 $0.63521 (18)$ $0.93764 (18)$ $0.54642 (18)$ $0.0354 (4)$ H36 0.5998 0.9532 0.4700 0.042^{*} N51 $0.22614 (19)$ $0.3177 (17)$ $0.0370 (4)$ H36 0.599 | C25 | -0.04382 (18) | 0.2171 (2) | 0.02057 (18) | 0.0437 (5) |
|---|------|---------------|--------------|---------------|------------|
| C26 $-0.02378 (17)$ $0.31482 (19)$ $0.09868 (16)$ $0.0334 (4)$ H26 -0.0449 0.3965 0.0745 0.040^* C41 $0.53859 (16)$ $0.58793 (18)$ $0.34023 (17)$ $0.0321 (4)$ H41A 0.5671 0.5174 0.3899 0.039^* C42 $0.56485 (16)$ $0.70599 (18)$ $0.40543 (17)$ $0.0327 (4)$ H42 0.5461 0.7823 0.3673 0.039^* C43 $0.61323 (16)$ $0.70778 (17)$ $0.51436 (16)$ $0.0313 (4)$ H43 0.6269 0.6293 0.5502 0.038^* C31 $0.64836 (15)$ $0.81551 (17)$ 0.5502 0.038^* C32 $0.70012 (17)$ $0.79671 (19)$ $0.69949 (17)$ $0.0342 (4)$ H32 0.7088 0.7147 0.7287 0.041^* C33 $0.73911 (18)$ $0.8947 (2)$ $0.77006 (18)$ 0.0408^* C34 $0.72614 (19)$ $1.0147 (2)$ $0.7288 (2)$ $0.0425 (4)$ H34 0.7552 1.0824 0.7767 0.051^* C35 0.6637 1.1180 0.5893 0.050^* C36 $0.63521 (18)$ $0.93764 (18)$ $0.54642 (18)$ $0.0342 (4)$ H36 0.5998 0.9532 0.4700 0.042^* N51 $0.26141 (19)$ $1.03197 (17)$ $-0.06212 (17)$ $0.0492 (5)$ C52 $0.2893 (19)$ $0.93764 (18)$ $0.0654 (17)$ $0.0376 (4)$ H36 0.5998 0.9522 0.4700 0.042^* N51 | H25 | -0.0782 | 0.2311 | -0.0565 | 0.052* |
| H26 -0.0449 0.3965 0.0745 0.040^* C41 $0.53859(16)$ $0.58793(18)$ $0.34023(17)$ $0.0321(4)$ H41A 0.5671 0.5174 0.3899 0.039^* C42 $0.56485(16)$ $0.70599(18)$ $0.40543(17)$ $0.0327(4)$ H42 0.5461 0.7823 0.3673 0.039^* C43 $0.61323(16)$ $0.70778(17)$ $0.51436(16)$ $0.0313(4)$ H43 0.6269 0.6293 0.5502 0.038^* C31 $0.64836(15)$ $0.81561(17)$ $0.58665(15)$ $0.0299(4)$ C32 $0.70012(17)$ $0.79671(19)$ $0.69949(17)$ $0.0342(4)$ H32 0.7088 0.7147 0.7287 0.041^* C33 $0.73911(18)$ $0.8947(2)$ $0.77006(18)$ $0.0401(4)$ H33 0.7747 0.8797 0.8466 0.048^* C34 $0.72614(19)$ $1.0147(2)$ $0.7288(2)$ $0.0425(4)$ H34 0.6637 1.1180 0.5893 0.050^* C35 0.6637 1.1180 0.5893 0.050^* C36 $0.63521(18)$ $0.93764(18)$ $0.54642(18)$ $0.0354(4)$ H36 0.5998 0.9532 0.4700 0.042^* N51 $0.26141(19)$ $0.3197(17)$ $-0.12234(17)$ $0.0376(4)$ H36 $0.5998(19)$ 0.9261 0.0774 0.044^* C53 $0.37557(16)$ $0.84493(16)$ $-0.1882(15)$ $0.0290(3)$ C54 $0.4063(2)$ $0.9557(2)$ -0.195 | C26 | -0.02378 (17) | 0.31482 (19) | 0.09868 (16) | 0.0334 (4) |
| C41 $0.53859 (16)$ $0.58793 (18)$ $0.34023 (17)$ $0.0321 (4)$ H41A 0.5671 0.5174 0.3899 $0.039*$ H41B 0.5860 0.5870 0.2681 $0.039*$ C42 $0.56485 (16)$ $0.70599 (18)$ $0.40543 (17)$ $0.0327 (4)$ H42 0.5461 0.7223 0.3673 $0.039*$ C43 $0.61323 (16)$ $0.70778 (17)$ $0.51436 (16)$ $0.0313 (4)$ H43 0.6269 0.6293 0.5502 $0.038*$ C31 $0.64836 (15)$ $0.81561 (17)$ $0.58665 (15)$ $0.0299 (4)$ C32 $0.70012 (17)$ $0.79671 (19)$ $0.69949 (17)$ $0.0342 (4)$ H32 0.7088 0.7147 0.7287 $0.041*$ C33 $0.73911 (18)$ $0.8947 (2)$ $0.77006 (18)$ $0.0401 (4)$ H33 0.7747 0.8797 0.8466 $0.048*$ C34 $0.72614 (19)$ $1.0147 (2)$ $0.7288 (2)$ $0.0425 (4)$ H34 0.7532 1.0824 0.7767 $0.051*$ C35 0.6371 1.180 0.5893 $0.050*$ C36 $0.63521 (18)$ $0.93764 (18)$ $0.54642 (18)$ $0.03354 (4)$ H36 0.5998 0.9532 0.4700 $0.442*$ N51 $0.26141 (19)$ $1.03197 (17)$ $-0.06212 (17)$ $0.0378 (4)$ H54 0.4972 0.7991 -0.1426 $0.045*$ C55 $0.4063 (2)$ $0.9557 (2)$ $-0.19576 (19)$ $0.0464 (5)$ H54 0.4972 0.95 | H26 | -0.0449 | 0.3965 | 0.0745 | 0.040* |
| H41A 0.5671 0.5174 0.3899 $0.039*$ H41B 0.5860 0.5870 0.2681 $0.039*$ C42 0.56485 (16) 0.70599 (18) 0.40543 (17) 0.0327 (4)H42 0.5461 0.7823 0.3673 $0.039*$ C43 0.61323 (16) 0.70778 (17) 0.51436 (16) 0.0313 (4)H43 0.6269 0.6293 0.5502 $0.038*$ C31 0.64836 (15) 0.81561 (17) 0.58665 (15) 0.0299 (4)C32 0.70012 (17) 0.79671 (19) 0.69949 (17) 0.0342 (4)H32 0.7088 0.7147 0.7287 $0.041*$ C33 0.73911 (18) 0.8947 (2) 0.77066 (18) 0.0401 (4)H33 0.7747 0.8797 0.84666 $0.0448*$ C34 0.72614 (19) 1.0147 (2) 0.7288 (2) 0.0425 (4)H34 0.7532 1.0824 0.7767 $0.051*$ C35 0.67351 (19) 1.03563 (19) 0.6173 (2) 0.0417 (4)H35 0.6637 1.1180 0.5893 $0.050*$ C36 0.63521 (18) 0.93764 (18) 0.54642 (18) 0.0354 (4)H36 0.5998 0.9532 0.4700 $0.042*$ N51 0.26141 (19) 1.03197 (17) -0.06212 (17) 0.0378 (4)H52 0.2473 0.9261 0.0774 $0.044*$ C53 0.37557 (16) 0.84493 (16) -0.12234 (17) 0.0378 (4)H54 0.4972 0 | C41 | 0.53859 (16) | 0.58793 (18) | 0.34023 (17) | 0.0321 (4) |
| H41B0.58600.58700.26810.039*C420.56485 (16)0.70599 (18)0.40543 (17)0.0327 (4)H420.54610.78230.36730.039*C430.61323 (16)0.70778 (17)0.51436 (16)0.0313 (4)H430.62690.62930.55020.038*C310.64836 (15)0.81561 (17)0.58665 (15)0.0299 (4)C320.70012 (17)0.79671 (19)0.69949 (17)0.0342 (4)H320.70880.71470.72870.041*C330.73911 (18)0.8947 (2)0.77066 (18)0.0401 (4)H330.77470.87970.84660.048*C340.72614 (19)1.0147 (2)0.7288 (2)0.0425 (4)H340.75321.08240.77670.051*C350.67351 (19)1.03563 (19)0.6173 (2)0.0417 (4)H350.66371.11800.58930.050*C360.63521 (18)0.93764 (18)0.54642 (18)0.0354 (4)H360.59980.95320.47000.042*N510.26141 (19)1.03197 (17)-0.06212 (17)0.0378 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.1882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.1957 (19)0.0464 (5) | H41A | 0.5671 | 0.5174 | 0.3899 | 0.039* |
| C42 $0.56485(16)$ $0.70599(18)$ $0.40543(17)$ $0.0327(4)$ H42 0.5461 0.7823 0.3673 $0.039*$ C43 $0.61323(16)$ $0.70778(17)$ $0.51436(16)$ $0.0313(4)$ H43 0.6269 0.6293 0.5502 $0.038*$ C31 $0.64836(15)$ $0.81561(17)$ $0.5865(15)$ $0.0299(4)$ C32 $0.70012(17)$ $0.79671(19)$ $0.69949(17)$ $0.0342(4)$ H32 0.7088 0.7147 0.7287 $0.041*$ C33 $0.73911(18)$ $0.8947(2)$ $0.77006(18)$ $0.0401(4)$ H33 0.7747 0.8797 0.8466 $0.048*$ C34 $0.72614(19)$ $1.0147(2)$ $0.7288(2)$ $0.0425(4)$ H34 0.7532 1.0824 0.7767 $0.051*$ C35 $0.67351(19)$ $1.03563(19)$ $0.6173(2)$ $0.0417(4)$ H35 0.6637 1.1180 0.5893 $0.050*$ C36 $0.63521(18)$ $0.93764(18)$ $0.54642(18)$ $0.0354(4)$ H36 0.5998 0.9532 0.4700 $0.042*$ N51 $0.26141(19)$ $1.03197(17)$ $-0.06212(17)$ $0.0492(5)$ C52 $0.28936(19)$ $0.93747(18)$ $0.00654(17)$ $0.0378(4)$ H54 0.4972 $0.9951(2)$ $-0.1182(15)$ $0.0290(3)$ C54 $0.43615(18)$ $0.85725(19)$ $-0.12234(17)$ $0.0378(4)$ H54 0.4972 $0.9951(2)$ $-0.1182(15)$ $0.0278(3)$ C55 $0.4063(2)$ 0.9 | H41B | 0.5860 | 0.5870 | 0.2681 | 0.039* |
| H42 0.5461 0.7823 0.3673 $0.039*$ C43 $0.61323 (16)$ $0.70778 (17)$ $0.51436 (16)$ $0.0313 (4)$ H43 0.6269 0.6293 0.5502 $0.038*$ C31 $0.64836 (15)$ $0.81561 (17)$ $0.58665 (15)$ $0.0299 (4)$ C32 $0.70012 (17)$ $0.79671 (19)$ $0.69949 (17)$ $0.0342 (4)$ H32 0.7088 0.7147 0.7287 $0.041*$ C33 $0.73911 (18)$ $0.8947 (2)$ $0.77006 (18)$ $0.0401 (4)$ H33 0.7747 0.8797 0.8466 $0.048*$ C34 $0.72614 (19)$ $1.0147 (2)$ $0.7288 (2)$ $0.0425 (4)$ H34 0.7532 1.0824 0.7767 $0.051*$ C35 $0.67351 (19)$ $1.03563 (19)$ $0.6173 (2)$ $0.0417 (4)$ H35 0.6637 1.1180 0.5893 $0.050*$ C36 $0.63521 (18)$ $0.93764 (18)$ $0.54642 (18)$ $0.0354 (4)$ H36 0.5998 0.9532 0.4700 $0.042*$ N51 $0.26141 (19)$ $1.03197 (17)$ $-0.06212 (17)$ $0.0492 (5)$ C52 $0.28936 (19)$ $0.93447 (18)$ $0.00654 (17)$ $0.0370 (4)$ H52 0.2473 0.9261 0.0774 $0.044*$ C53 $0.37557 (16)$ $0.85725 (19)$ $-0.12234 (17)$ $0.0378 (4)$ H54 0.4972 0.7991 -0.1426 $0.045*$ C55 $0.4063 (2)$ $0.9557 (2)$ $-0.1618 (2)$ $0.0480 (5)$ H55 0.44 | C42 | 0.56485 (16) | 0.70599 (18) | 0.40543 (17) | 0.0327 (4) |
| C43 $0.61323 (16)$ $0.70778 (17)$ $0.51436 (16)$ $0.0313 (4)$ H43 0.6269 0.6293 0.5502 $0.038*$ C31 $0.64836 (15)$ $0.81561 (17)$ $0.58665 (15)$ $0.0299 (4)$ C32 $0.70012 (17)$ $0.79671 (19)$ $0.69949 (17)$ $0.0342 (4)$ H32 0.7088 0.7147 0.7287 $0.041*$ C33 $0.73911 (18)$ $0.8947 (2)$ $0.7006 (18)$ $0.0401 (4)$ H33 0.7747 0.8797 0.8466 $0.048*$ C34 $0.72614 (19)$ $1.0147 (2)$ $0.7288 (2)$ $0.0425 (4)$ H34 0.7532 1.0824 0.7767 $0.051*$ C35 $0.67351 (19)$ $1.03563 (19)$ $0.6173 (2)$ $0.0417 (4)$ H35 0.6637 1.1180 0.5893 $0.050*$ C36 $0.63521 (18)$ $0.93764 (18)$ $0.54642 (18)$ $0.0354 (4)$ H36 0.5998 0.9532 0.4700 $0.042*$ N51 $0.26141 (19)$ $1.03197 (17)$ $-0.06212 (17)$ $0.0492 (5)$ C52 $0.28936 (19)$ $0.93447 (18)$ $0.00654 (17)$ $0.0370 (4)$ H52 0.2473 0.9261 0.0774 $0.044*$ C53 $0.37557 (16)$ $0.84493 (16)$ $-0.1182 (15)$ $0.0290 (3)$ C54 $0.43615 (18)$ $0.85725 (19)$ $-0.1228 (17)$ $0.0378 (4)$ H54 0.4972 0.7991 -0.1426 $0.045*$ C55 $0.4063 (2)$ $0.9557 (2)$ $-0.19576 (19)$ $0.0464 (5)$ <tr< td=""><td>H42</td><td>0.5461</td><td>0.7823</td><td>0.3673</td><td>0.039*</td></tr<> | H42 | 0.5461 | 0.7823 | 0.3673 | 0.039* |
| H430.62690.62930.55020.038*C310.64836 (15)0.81561 (17)0.58665 (15)0.0299 (4)C320.70012 (17)0.79671 (19)0.69949 (17)0.0342 (4)H320.70880.71470.72870.041*C330.73911 (18)0.8947 (2)0.77006 (18)0.0401 (4)H330.77470.87970.84660.048*C340.72614 (19)1.0147 (2)0.7288 (2)0.0425 (4)H340.75321.08240.77670.051*C350.67351 (19)1.03563 (19)0.6173 (2)0.0417 (4)H350.66371.11800.58930.050*C360.63521 (18)0.93764 (18)0.54642 (18)0.0354 (4)H360.59980.95320.47000.042*N510.26141 (19)1.03197 (17)-0.06212 (17)0.0492 (5)C520.28936 (19)0.93447 (18)0.00654 (17)0.0370 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.11882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4463 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0488 (5)H560.30151.1076-0.21250.058* <t< td=""><td>C43</td><td>0.61323 (16)</td><td>0.70778 (17)</td><td>0.51436 (16)</td><td>0.0313 (4)</td></t<> | C43 | 0.61323 (16) | 0.70778 (17) | 0.51436 (16) | 0.0313 (4) |
| C310.64836 (15)0.81561 (17)0.58665 (15)0.0299 (4)C320.70012 (17)0.79671 (19)0.69949 (17)0.0342 (4)H320.70880.71470.72870.041*C330.73911 (18)0.8947 (2)0.77006 (18)0.0401 (4)H330.77470.87970.84660.048*C340.72614 (19)1.0147 (2)0.7288 (2)0.0425 (4)H340.75321.08240.77670.051*C350.67351 (19)1.03563 (19)0.6173 (2)0.0417 (4)H350.66371.11800.58930.050*C360.63521 (18)0.93764 (18)0.54642 (18)0.0354 (4)H360.59980.95320.47000.042*N510.26141 (19)1.03197 (17)-0.06212 (17)0.0492 (5)C520.28936 (19)0.93447 (18)0.00654 (17)0.0370 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.1882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.66383 (15)0 | H43 | 0.6269 | 0.6293 | 0.5502 | 0.038* |
| C320.70012 (17)0.79671 (19)0.69949 (17)0.0342 (4)H320.70880.71470.72870.041*C330.73911 (18)0.8947 (2)0.77006 (18)0.0401 (4)H330.77470.87970.84660.048*C340.72614 (19)1.0147 (2)0.7288 (2)0.0425 (4)H340.75321.08240.77670.051*C350.67351 (19)1.03563 (19)0.6173 (2)0.0417 (4)H350.66371.11800.58930.050*C360.63521 (18)0.93764 (18)0.54642 (18)0.0354 (4)H360.59980.95320.47000.042*N510.26141 (19)1.03197 (17)-0.06212 (17)0.0492 (5)C520.28936 (19)0.93447 (18)0.00654 (17)0.0370 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.11234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0 | C31 | 0.64836 (15) | 0.81561 (17) | 0.58665 (15) | 0.0299 (4) |
| H320.70880.71470.72870.041*C330.73911 (18)0.8947 (2)0.77006 (18)0.0401 (4)H330.77470.87970.84660.048*C340.72614 (19)1.0147 (2)0.7288 (2)0.0425 (4)H340.75321.08240.77670.051*C350.67351 (19)1.03563 (19)0.6173 (2)0.0417 (4)H350.66371.11800.58930.050*C360.63521 (18)0.93764 (18)0.54642 (18)0.0354 (4)H360.59980.95320.47000.042*N510.26141 (19)1.03197 (17)-0.06212 (17)0.0492 (5)C520.28936 (19)0.93447 (18)0.00654 (17)0.0370 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.01882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13) | C32 | 0.70012 (17) | 0.79671 (19) | 0.69949 (17) | 0.0342 (4) |
| C330.73911 (18)0.8947 (2)0.77006 (18)0.0401 (4)H330.77470.87970.84660.048*C340.72614 (19)1.0147 (2)0.7288 (2)0.0425 (4)H340.75321.08240.77670.051*C350.67351 (19)1.03563 (19)0.6173 (2)0.0417 (4)H350.66371.11800.58930.050*C360.63521 (18)0.93764 (18)0.54642 (18)0.0354 (4)H360.59980.95320.47000.042*N510.26141 (19)1.03197 (17)-0.06212 (17)0.0492 (5)C520.28936 (19)0.93447 (18)0.00654 (17)0.0370 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.01882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06333 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | H32 | 0.7088 | 0.7147 | 0.7287 | 0.041* |
| H330.77470.87970.84660.048*C340.72614 (19)1.0147 (2)0.7288 (2)0.0425 (4)H340.75321.08240.77670.051*C350.67351 (19)1.03563 (19)0.6173 (2)0.0417 (4)H350.66371.11800.58930.050*C360.63521 (18)0.93764 (18)0.54642 (18)0.0354 (4)H360.59980.95320.47000.042*N510.26141 (19)1.03197 (17)-0.06212 (17)0.0492 (5)C520.28936 (19)0.93447 (18)0.00654 (17)0.0370 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.1882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | C33 | 0.73911 (18) | 0.8947 (2) | 0.77006 (18) | 0.0401 (4) |
| C340.72614 (19)1.0147 (2)0.7288 (2)0.0425 (4)H340.75321.08240.77670.051*C350.67351 (19)1.03563 (19)0.6173 (2)0.0417 (4)H350.66371.11800.58930.050*C360.63521 (18)0.93764 (18)0.54642 (18)0.0354 (4)H360.59980.95320.47000.042*N510.26141 (19)1.03197 (17)-0.06212 (17)0.0492 (5)C520.28936 (19)0.93447 (18)0.00654 (17)0.0370 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.01882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | H33 | 0.7747 | 0.8797 | 0.8466 | 0.048* |
| H340.75321.08240.77670.051*C350.67351 (19)1.03563 (19)0.6173 (2)0.0417 (4)H350.66371.11800.58930.050*C360.63521 (18)0.93764 (18)0.54642 (18)0.0354 (4)H360.59980.95320.47000.042*N510.26141 (19)1.03197 (17)-0.06212 (17)0.0492 (5)C520.28936 (19)0.93447 (18)0.00654 (17)0.0370 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.01882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.31051.1076-0.21250.058*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.4047 (3) | C34 | 0.72614 (19) | 1.0147 (2) | 0.7288 (2) | 0.0425 (4) |
| C350.67351 (19)1.03563 (19)0.6173 (2)0.0417 (4)H350.66371.11800.58930.050*C360.63521 (18)0.93764 (18)0.54642 (18)0.0354 (4)H360.59980.95320.47000.042*N510.26141 (19)1.03197 (17)-0.06212 (17)0.0492 (5)C520.28936 (19)0.93447 (18)0.00654 (17)0.0370 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.1882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | H34 | 0.7532 | 1.0824 | 0.7767 | 0.051* |
| H350.66371.11800.58930.050*C360.63521 (18)0.93764 (18)0.54642 (18)0.0354 (4)H360.59980.95320.47000.042*N510.26141 (19)1.03197 (17)-0.06212 (17)0.0492 (5)C520.28936 (19)0.93447 (18)0.00654 (17)0.0370 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.01882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | C35 | 0.67351 (19) | 1.03563 (19) | 0.6173 (2) | 0.0417 (4) |
| C36 $0.63521(18)$ $0.93764(18)$ $0.54642(18)$ $0.0354(4)$ H36 0.5998 0.9532 0.4700 0.042^* N51 $0.26141(19)$ $1.03197(17)$ $-0.06212(17)$ $0.0492(5)$ C52 $0.28936(19)$ $0.93447(18)$ $0.00654(17)$ $0.0370(4)$ H52 0.2473 0.9261 0.0774 0.044^* C53 $0.37557(16)$ $0.84493(16)$ $-0.01882(15)$ $0.0290(3)$ C54 $0.43615(18)$ $0.85725(19)$ $-0.12234(17)$ $0.0378(4)$ H54 0.4972 0.7991 -0.1426 0.045^* C55 $0.4063(2)$ $0.9557(2)$ $-0.19576(19)$ $0.0464(5)$ H55 0.4447 0.9650 -0.2685 0.056^* C56 $0.3202(2)$ $1.0397(2)$ $-0.1618(2)$ $0.0480(5)$ H56 0.3015 1.1076 -0.2125 0.058^* C57 $0.40362(16)$ $0.73877(16)$ $0.06383(15)$ $0.0278(3)$ O51 $0.33722(13)$ $0.73420(12)$ $0.15347(11)$ $0.0360(3)$ O52 $0.48540(13)$ $0.66376(13)$ $0.04174(13)$ $0.0407(3)$ | H35 | 0.6637 | 1.1180 | 0.5893 | 0.050* |
| H36 0.5998 0.9532 0.4700 $0.042*$ N51 $0.26141(19)$ $1.03197(17)$ $-0.06212(17)$ $0.0492(5)$ C52 $0.28936(19)$ $0.93447(18)$ $0.00654(17)$ $0.0370(4)$ H52 0.2473 0.9261 0.0774 $0.044*$ C53 $0.37557(16)$ $0.84493(16)$ $-0.01882(15)$ $0.0290(3)$ C54 $0.43615(18)$ $0.85725(19)$ $-0.12234(17)$ $0.0378(4)$ H54 0.4972 0.7991 -0.1426 $0.045*$ C55 $0.4063(2)$ $0.9557(2)$ $-0.19576(19)$ $0.0464(5)$ H55 0.4447 0.9650 -0.2685 $0.056*$ C56 $0.3202(2)$ $1.0397(2)$ $-0.1618(2)$ $0.0480(5)$ H56 0.3015 1.1076 -0.2125 $0.058*$ C57 $0.40362(16)$ $0.73877(16)$ $0.06383(15)$ $0.0278(3)$ O51 $0.33722(13)$ $0.73420(12)$ $0.15347(11)$ $0.0360(3)$ O52 $0.48540(13)$ $0.66376(13)$ $0.04174(13)$ $0.0407(3)$ | C36 | 0.63521 (18) | 0.93764 (18) | 0.54642 (18) | 0.0354 (4) |
| N510.26141 (19)1.03197 (17)-0.06212 (17)0.0492 (5)C520.28936 (19)0.93447 (18)0.00654 (17)0.0370 (4)H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.01882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | H36 | 0.5998 | 0.9532 | 0.4700 | 0.042* |
| C52 $0.28936(19)$ $0.93447(18)$ $0.00654(17)$ $0.0370(4)$ H52 0.2473 0.9261 0.0774 $0.044*$ C53 $0.37557(16)$ $0.84493(16)$ $-0.01882(15)$ $0.0290(3)$ C54 $0.43615(18)$ $0.85725(19)$ $-0.12234(17)$ $0.0378(4)$ H54 0.4972 0.7991 -0.1426 $0.045*$ C55 $0.4063(2)$ $0.9557(2)$ $-0.19576(19)$ $0.0464(5)$ H55 0.4447 0.9650 -0.2685 $0.056*$ C56 $0.3202(2)$ $1.0397(2)$ $-0.1618(2)$ $0.0480(5)$ H56 0.3015 1.1076 -0.2125 $0.058*$ C57 $0.40362(16)$ $0.73877(16)$ $0.06383(15)$ $0.0278(3)$ O51 $0.33722(13)$ $0.73420(12)$ $0.15347(11)$ $0.0360(3)$ O52 $0.48540(13)$ $0.66376(13)$ $0.04174(13)$ $0.0407(3)$ | N51 | 0.26141 (19) | 1.03197 (17) | -0.06212 (17) | 0.0492 (5) |
| H520.24730.92610.07740.044*C530.37557 (16)0.84493 (16)-0.01882 (15)0.0290 (3)C540.43615 (18)0.85725 (19)-0.12234 (17)0.0378 (4)H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | C52 | 0.28936 (19) | 0.93447 (18) | 0.00654 (17) | 0.0370 (4) |
| C53 $0.37557 (16)$ $0.84493 (16)$ $-0.01882 (15)$ $0.0290 (3)$ C54 $0.43615 (18)$ $0.85725 (19)$ $-0.12234 (17)$ $0.0378 (4)$ H54 0.4972 0.7991 -0.1426 0.045^* C55 $0.4063 (2)$ $0.9557 (2)$ $-0.19576 (19)$ $0.0464 (5)$ H55 0.4447 0.9650 -0.2685 0.056^* C56 $0.3202 (2)$ $1.0397 (2)$ $-0.1618 (2)$ $0.0480 (5)$ H56 0.3015 1.1076 -0.2125 0.058^* C57 $0.40362 (16)$ $0.73877 (16)$ $0.06383 (15)$ $0.0278 (3)$ O51 $0.33722 (13)$ $0.73420 (12)$ $0.15347 (11)$ $0.0360 (3)$ O52 $0.48540 (13)$ $0.66376 (13)$ $0.04174 (13)$ $0.0407 (3)$ | H52 | 0.2473 | 0.9261 | 0.0774 | 0.044* |
| C54 $0.43615(18)$ $0.85725(19)$ $-0.12234(17)$ $0.0378(4)$ H54 0.4972 0.7991 -0.1426 $0.045*$ C55 $0.4063(2)$ $0.9557(2)$ $-0.19576(19)$ $0.0464(5)$ H55 0.4447 0.9650 -0.2685 $0.056*$ C56 $0.3202(2)$ $1.0397(2)$ $-0.1618(2)$ $0.0480(5)$ H56 0.3015 1.1076 -0.2125 $0.058*$ C57 $0.40362(16)$ $0.73877(16)$ $0.66383(15)$ $0.0278(3)$ O51 $0.33722(13)$ $0.73420(12)$ $0.15347(11)$ $0.0360(3)$ O52 $0.48540(13)$ $0.66376(13)$ $0.04174(13)$ $0.0407(3)$ | C53 | 0.37557 (16) | 0.84493 (16) | -0.01882 (15) | 0.0290 (3) |
| H540.49720.7991-0.14260.045*C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | C54 | 0.43615 (18) | 0.85725 (19) | -0.12234 (17) | 0.0378 (4) |
| C550.4063 (2)0.9557 (2)-0.19576 (19)0.0464 (5)H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | H54 | 0.4972 | 0.7991 | -0.1426 | 0.045* |
| H550.44470.9650-0.26850.056*C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | C55 | 0.4063 (2) | 0.9557 (2) | -0.19576 (19) | 0.0464 (5) |
| C560.3202 (2)1.0397 (2)-0.1618 (2)0.0480 (5)H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | H55 | 0.4447 | 0.9650 | -0.2685 | 0.056* |
| H560.30151.1076-0.21250.058*C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | C56 | 0.3202 (2) | 1.0397 (2) | -0.1618 (2) | 0.0480 (5) |
| C570.40362 (16)0.73877 (16)0.06383 (15)0.0278 (3)O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | H56 | 0.3015 | 1.1076 | -0.2125 | 0.058* |
| O510.33722 (13)0.73420 (12)0.15347 (11)0.0360 (3)O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | C57 | 0.40362 (16) | 0.73877 (16) | 0.06383 (15) | 0.0278 (3) |
| O520.48540 (13)0.66376 (13)0.04174 (13)0.0407 (3) | O51 | 0.33722 (13) | 0.73420 (12) | 0.15347 (11) | 0.0360 (3) |
| | 052 | 0.48540 (13) | 0.66376 (13) | 0.04174 (13) | 0.0407 (3) |

Atomic displacement parameters $(Å^2)$

| | U^{11} | U^{22} | U^{33} | U^{12} | U^{13} | U^{23} |
|-----|------------|------------|------------|-------------|------------|-------------|
| N1 | 0.0236 (6) | 0.0273 (7) | 0.0243 (7) | -0.0025 (5) | 0.0046 (5) | -0.0024 (5) |
| C2 | 0.0252 (7) | 0.0279 (8) | 0.0255 (8) | -0.0009 (6) | 0.0040 (6) | -0.0033 (6) |
| C3 | 0.0277 (8) | 0.0318 (8) | 0.0212 (8) | -0.0038 (6) | 0.0037 (6) | -0.0027 (6) |
| N4 | 0.0222 (6) | 0.0288 (7) | 0.0222 (7) | -0.0022 (5) | 0.0019 (5) | 0.0008 (5) |
| C5 | 0.0259 (8) | 0.0296 (8) | 0.0277 (8) | -0.0003 (6) | 0.0053 (6) | -0.0025 (7) |
| C6 | 0.0262 (8) | 0.0316 (8) | 0.0245 (8) | -0.0026 (6) | 0.0046 (6) | -0.0039 (6) |
| C1 | 0.0230 (7) | 0.0257 (7) | 0.0262 (8) | -0.0001 (6) | 0.0008 (6) | 0.0006 (6) |
| C11 | 0.0247 (7) | 0.0235 (7) | 0.0274 (8) | -0.0033 (6) | 0.0028 (6) | -0.0037 (6) |
| C12 | 0.0245 (7) | 0.0332 (8) | 0.0327 (9) | -0.0001 (6) | 0.0000 (6) | -0.0040 (7) |
| | | | | | | |

| C13 | 0.0271 (8) | 0.0378 (9) | 0.0432 (10) | -0.0044 (7) | 0.0089 (7) | -0.0084 (8) |
|-----|-------------|-------------|-------------|--------------|--------------|--------------|
| C14 | 0.0405 (9) | 0.0319 (8) | 0.0323 (9) | -0.0098 (7) | 0.0141 (7) | -0.0047 (7) |
| F14 | 0.0582 (8) | 0.0568 (8) | 0.0419 (7) | -0.0109 (6) | 0.0256 (6) | -0.0005 (6) |
| C15 | 0.0407 (9) | 0.0307 (8) | 0.0294 (9) | -0.0036 (7) | 0.0023 (7) | 0.0016 (7) |
| C16 | 0.0266 (8) | 0.0298 (8) | 0.0319 (9) | 0.0001 (6) | 0.0027 (6) | 0.0002 (7) |
| C21 | 0.0221 (7) | 0.0305 (8) | 0.0270 (8) | -0.0039 (6) | 0.0020 (6) | -0.0027 (7) |
| C22 | 0.0351 (9) | 0.0322 (9) | 0.0364 (10) | -0.0002 (7) | 0.0018 (7) | -0.0027 (7) |
| C23 | 0.0410 (10) | 0.0352 (10) | 0.0550 (13) | -0.0023 (8) | 0.0095 (9) | -0.0120 (9) |
| C24 | 0.0364 (10) | 0.0527 (12) | 0.0479 (12) | -0.0120 (9) | 0.0108 (9) | -0.0264 (10) |
| F24 | 0.0647 (9) | 0.0721 (10) | 0.0744 (10) | -0.0163 (8) | 0.0103 (8) | -0.0485 (8) |
| C25 | 0.0344 (9) | 0.0675 (14) | 0.0292 (9) | -0.0095 (9) | 0.0023 (8) | -0.0119 (9) |
| C26 | 0.0289 (8) | 0.0439 (10) | 0.0278 (9) | -0.0041 (7) | 0.0029 (7) | -0.0008 (7) |
| C41 | 0.0227 (7) | 0.0409 (9) | 0.0327 (9) | -0.0019 (7) | 0.0003 (6) | 0.0000(7) |
| C42 | 0.0262 (8) | 0.0358 (9) | 0.0359 (10) | -0.0046 (7) | 0.0004 (7) | 0.0046 (8) |
| C43 | 0.0276 (8) | 0.0323 (8) | 0.0341 (9) | 0.0022 (7) | 0.0022 (7) | 0.0020 (7) |
| C31 | 0.0226 (7) | 0.0364 (9) | 0.0309 (9) | 0.0012 (6) | 0.0026 (6) | 0.0005 (7) |
| C32 | 0.0300 (8) | 0.0392 (9) | 0.0334 (9) | 0.0048 (7) | 0.0013 (7) | 0.0025 (8) |
| C33 | 0.0338 (9) | 0.0545 (12) | 0.0316 (9) | 0.0006 (8) | -0.0019 (7) | -0.0019 (9) |
| C34 | 0.0359 (9) | 0.0458 (11) | 0.0458 (11) | -0.0066 (8) | 0.0020 (8) | -0.0105 (9) |
| C35 | 0.0393 (10) | 0.0342 (9) | 0.0516 (12) | -0.0031 (8) | 0.0031 (9) | 0.0008 (9) |
| C36 | 0.0331 (9) | 0.0392 (9) | 0.0335 (9) | 0.0003 (7) | -0.0012 (7) | 0.0046 (8) |
| N51 | 0.0569 (11) | 0.0434 (10) | 0.0469 (11) | 0.0060 (9) | -0.0031 (9) | 0.0113 (8) |
| C52 | 0.0416 (10) | 0.0388 (10) | 0.0309 (9) | 0.0023 (8) | 0.0040 (7) | 0.0052 (8) |
| C53 | 0.0299 (8) | 0.0299 (8) | 0.0271 (8) | -0.0083 (6) | -0.0002 (6) | -0.0008 (7) |
| C54 | 0.0387 (9) | 0.0413 (10) | 0.0342 (10) | -0.0121 (8) | 0.0091 (8) | -0.0037 (8) |
| C55 | 0.0548 (12) | 0.0545 (12) | 0.0302 (10) | -0.0224 (10) | 0.0054 (9) | 0.0076 (9) |
| C56 | 0.0563 (13) | 0.0447 (11) | 0.0418 (12) | -0.0128 (10) | -0.0097 (10) | 0.0161 (9) |
| C57 | 0.0280 (7) | 0.0275 (8) | 0.0279 (8) | -0.0049 (6) | -0.0006 (6) | -0.0020 (6) |
| O51 | 0.0428 (7) | 0.0361 (6) | 0.0298 (6) | 0.0054 (6) | 0.0086 (5) | 0.0067 (5) |
| O52 | 0.0382 (7) | 0.0389 (7) | 0.0457 (8) | 0.0061 (6) | 0.0082 (6) | -0.0006 (6) |
| | | | | | | |

Geometric parameters (Å, °)

| N1—C6 | 1.463 (2) | C24—F24 | 1.355 (2) |
|--------|-----------|----------|-----------|
| N1—C2 | 1.469 (2) | C24—C25 | 1.368 (3) |
| N1-C1 | 1.479 (2) | C25—C26 | 1.391 (3) |
| С2—С3 | 1.512 (2) | C25—H25 | 0.9500 |
| C2—H2A | 0.9900 | C26—H26 | 0.9500 |
| C2—H2B | 0.9900 | C41—C42 | 1.497 (3) |
| C3—N4 | 1.491 (2) | C41—H41A | 0.9900 |
| С3—НЗА | 0.9900 | C41—H41B | 0.9900 |
| С3—Н3В | 0.9900 | C42—C43 | 1.324 (3) |
| N4—C5 | 1.489 (2) | C42—H42 | 0.9500 |
| N4-C41 | 1.501 (2) | C43—C31 | 1.467 (3) |
| N4—H4 | 0.9807 | C43—H43 | 0.9500 |
| C5—C6 | 1.516 (2) | C31—C32 | 1.393 (3) |
| C5—H5A | 0.9900 | C31—C36 | 1.402 (3) |
| С5—Н5В | 0.9900 | C32—C33 | 1.384 (3) |
| | | | |

| | 0.0000 | C22 U22 | 0.0500 |
|----------------------------------|----------------------|--------------------------------------|-------------|
| | 0.9900 | C32—H32 | 0.9500 |
| C6—H6B | 0.9900 | C33—C34 | 1.385 (3) |
| C1—C21 | 1.515 (2) | С33—Н33 | 0.9500 |
| C1—C11 | 1.524 (2) | C34—C35 | 1.386 (3) |
| C1—H1 | 1.0000 | С34—Н34 | 0.9500 |
| C11—C16 | 1.388 (2) | C35—C36 | 1.384 (3) |
| C11—C12 | 1.396 (2) | С35—Н35 | 0.9500 |
| C12—C13 | 1.390 (3) | С36—Н36 | 0.9500 |
| C12—H12 | 0.9500 | N51—C56 | 1.332 (3) |
| C13—C14 | 1.364 (3) | N51—C52 | 1.339 (3) |
| С13—Н13 | 0.9500 | C52—C53 | 1.386 (3) |
| C14—F14 | 1 370 (2) | C52—H52 | 0.9500 |
| C14 $C15$ | 1.373(2) | C_{53} C_{54} | 1.384(2) |
| C15 C16 | 1.373(3) 1 302(3) | $C_{55} = C_{57}$ | 1.504(2) |
| C15_U15 | 0.0500 | C54 - C55 | 1.300(2) |
| | 0.9300 | C54_C55 | 1.383 (3) |
| | 0.9500 | C34—H54 | 0.9500 |
| C21—C26 | 1.394 (2) | C55—C56 | 1.371 (4) |
| C21—C22 | 1.397 (3) | С55—Н55 | 0.9500 |
| C22—C23 | 1.387 (3) | С56—Н56 | 0.9500 |
| C22—H22 | 0.9500 | C57—O52 | 1.237 (2) |
| C23—C24 | 1.371 (4) | C57—O51 | 1.278 (2) |
| С23—Н23 | 0.9500 | | |
| | | | |
| C6—N1—C2 | 108.19 (12) | C21—C22—H22 | 119.7 |
| C6—N1—C1 | 113.38 (12) | C24—C23—C22 | 118.8 (2) |
| C2—N1—C1 | 109.77 (12) | С24—С23—Н23 | 120.6 |
| N1—C2—C3 | 111.08 (13) | С22—С23—Н23 | 120.6 |
| N1—C2—H2A | 109.4 | F24—C24—C25 | 119.0 (2) |
| C3—C2—H2A | 109.4 | F24—C24—C23 | 118.2 (2) |
| N1—C2—H2B | 109.4 | C25—C24—C23 | 122.72 (18) |
| $C_3 = C_2 = H_2 B$ | 109.4 | C_{24} C_{25} C_{26} | 118 30 (19) |
| $H_2 \Delta (C_2) = H_2 B$ | 109.4 | $C_{24} = C_{25} = C_{20}$ | 120.8 |
| $M_{12} = C_2 = M_{12} = M_{12}$ | 100.0 110.27(12) | $C_{24} = C_{25} = H_{25}$ | 120.8 |
| N4 C2 U2A | 100.6 | $C_{20} = C_{23} = M_{23}$ | 120.0 |
| N4-C3-H3A | 109.0 | $C_{23} = C_{20} = C_{21}$ | 121.05 (19) |
| $C_2 - C_3 - H_3 A$ | 109.6 | C_{25} — C_{26} —H ₂₆ | 119.5 |
| N4—C3—H3B | 109.6 | C21—C26—H26 | 119.5 |
| С2—С3—Н3В | 109.6 | C42—C41—N4 | 112.91 (15) |
| НЗА—СЗ—НЗВ | 108.1 | C42—C41—H41A | 109.0 |
| C5—N4—C3 | 109.54 (12) | N4—C41—H41A | 109.0 |
| C5—N4—C41 | 109.36 (13) | C42—C41—H41B | 109.0 |
| C3—N4—C41 | 112.14 (13) | N4—C41—H41B | 109.0 |
| C5—N4—H4 | 108.3 | H41A—C41—H41B | 107.8 |
| C3—N4—H4 | 107.8 | C43—C42—C41 | 122.33 (17) |
| C41—N4—H4 | 109.6 | C43—C42—H42 | 118.8 |
| N4—C5—C6 | 110.88 (13) | C41—C42—H42 | 118.8 |
| N4—C5—H5A | 109.5 | C42—C43—C31 | 128.20 (17) |
| C6—C5—H5A | 109.5 | C42—C43—H43 | 115.9 |
| N4—C5—H5B | 109.5 | C31—C43—H43 | 115.9 |
| | | | |

| C6—C5—H5B | 109.5 | C32—C31—C36 | 118.03 (17) |
|---------------------------------|--------------------------|---|--------------------------|
| H5A—C5—H5B | 108.1 | C32—C31—C43 | 118.94 (16) |
| N1—C6—C5 | 109.75 (13) | C36—C31—C43 | 123.00 (16) |
| N1—C6—H6A | 109.7 | C33—C32—C31 | 121.50 (18) |
| С5—С6—Н6А | 109 7 | C33—C32—H32 | 1193 |
| N1—C6—H6B | 109.7 | C_{31} C_{32} H_{32} | 119.3 |
| C5 C6 H6B | 109.7 | C_{32} C_{33} C_{34} | 119.5 |
| | 109.7 | $C_{32} = C_{33} = C_{34}$ | 119.79 (19) |
| | 100.2 | C34 C32 H32 | 120.1 |
| NI - CI - C2I | 112.03 (13) | C34—C33—H33 | 120.1 |
| NI-CI-CII | 110.05 (13) | C33—C34—C35 | 119.62 (19) |
| C21—C1—C11 | 110.90 (13) | C33—C34—H34 | 120.2 |
| N1—C1—H1 | 107.9 | С35—С34—Н34 | 120.2 |
| C21—C1—H1 | 107.9 | C36—C35—C34 | 120.61 (19) |
| C11—C1—H1 | 107.9 | С36—С35—Н35 | 119.7 |
| C16—C11—C12 | 118.57 (16) | С34—С35—Н35 | 119.7 |
| C16—C11—C1 | 121.68 (14) | C35—C36—C31 | 120.44 (18) |
| C12—C11—C1 | 119.65 (15) | С35—С36—Н36 | 119.8 |
| C13—C12—C11 | 121.12 (16) | C31—C36—H36 | 119.8 |
| C13—C12—H12 | 119.4 | C56—N51—C52 | 116 3 (2) |
| $C_{11} - C_{12} - H_{12}$ | 119.4 | N51-C52-C53 | 124.38(19) |
| C14 $C13$ $C12$ | 117.95 (17) | N51 C52 H52 | 117.8 |
| $C_{14} = C_{13} = C_{12}$ | 117.95 (17) | 131 - 032 - 1152 | 117.8 |
| С14—С13—Н13 | 121.0 | С53—С52—П52 | 117.6 |
| C12—C13—H13 | 121.0 | C54—C53—C52 | 117.59 (17) |
| C13—C14—F14 | 118.32 (17) | C54—C53—C57 | 121.05 (17) |
| C13—C14—C15 | 123.42 (17) | C52—C53—C57 | 121.35 (16) |
| F14—C14—C15 | 118.25 (18) | C55—C54—C53 | 118.8 (2) |
| C14—C15—C16 | 117.93 (17) | С55—С54—Н54 | 120.6 |
| C14—C15—H15 | 121.0 | С53—С54—Н54 | 120.6 |
| C16—C15—H15 | 121.0 | C56—C55—C54 | 118.9 (2) |
| C11—C16—C15 | 120.99 (16) | С56—С55—Н55 | 120.6 |
| C11—C16—H16 | 119.5 | С54—С55—Н55 | 120.6 |
| C15—C16—H16 | 119.5 | N51—C56—C55 | 124.01 (19) |
| C26—C21—C22 | 118.54 (16) | N51—C56—H56 | 118.0 |
| $C_{26} = C_{21} = C_{1}$ | 119 76 (16) | С55—С56—Н56 | 118.0 |
| C_{22} C_{21} C_{1} | 121.69 (15) | 052 - 057 - 051 | 125.05(17) |
| C_{22} C_{21} C_{1} | 121.09(19) 120.62(19) | 052 057 051 | 129.05(17) 110.87(16) |
| $C_{23} = C_{22} = C_{21}$ | 110.7 | 051 C57 C53 | 115.07(10) |
| C23—C22—H22 | 119.7 | 051-057-055 | 113.07 (13) |
| C6—N1—C2—C3 | -61.39 (16) | C21—C22—C23—C24 | 0.3 (3) |
| C1—N1—C2—C3 | 174.41 (13) | C22—C23—C24—F24 | -178.91 (18) |
| N1-C2-C3-N4 | 58 60 (17) | C^{22} C^{23} C^{24} C^{25} | 0.0(3) |
| $C_2 - C_3 - N_4 - C_5$ | -54.81(17) | F_{24} C_{24} C_{25} C_{26} | 17874(18) |
| $C_2 C_3 N_4 C_4 1$ | -176 42 (14) | $C_{23}^{23} = C_{24}^{24} = C_{25}^{25} = C_{26}^{26}$ | -0.1(3) |
| $C_2 = C_3 = 14 = C_{11}$ | 55.00(17) | $C_{23} = C_{24} = C_{23} = C_{20} = C_{20}$ | 0.1(3) |
| $C_{1} = 104 - C_{2} - C_{0}$ | 33.77(17) | $C_{24} = C_{23} = C_{20} = C_{21}$ | 0.1(3) |
| $C_{1} = N_{1} = C_{2} = C_{0}$ | 1/9.20 (14) | 122 - 121 - 120 - 120 | 0.1 (3) |
| C2—N1—C6—C5 | 61.43 (16) | C1—C21—C26—C25 | -179.35 (16) |
| C1—N1—C6—C5 | -1/6.56 (14) | C5—N4—C41—C42 | -177.30 (14) |
| N4—C5—C6—N1 | -60.11 (17) | C3—N4—C41—C42 | -55.58 (19) |

| C6—N1—C1—C21 | 44.09 (18) | N4—C41—C42—C43 | 118.27 (19) |
|-----------------|--------------|-----------------|--------------|
| C6—N1—C1—C11 | 167.96 (13) | C41—C42—C43—C31 | 177.23 (16) |
| C2-N1-C1-C11 | -70.92 (16) | C42—C43—C31—C32 | -178.93 (18) |
| C2-N1-C1-C21 | 165.22 (13) | C42—C43—C31—C36 | -0.6 (3) |
| N1-C1-C11-C16 | -33.4 (2) | C36—C31—C32—C33 | -0.8 (3) |
| C21—C1—C11—C16 | 91.08 (18) | C43—C31—C32—C33 | 177.64 (17) |
| N1-C1-C11-C12 | 150.17 (15) | C31—C32—C33—C34 | 0.4 (3) |
| C21—C1—C11—C12 | -85.31 (18) | C32—C33—C34—C35 | 0.4 (3) |
| C16—C11—C12—C13 | -0.3 (2) | C33—C34—C35—C36 | -0.8 (3) |
| C1-C11-C12-C13 | 176.25 (15) | C34—C35—C36—C31 | 0.4 (3) |
| C11—C12—C13—C14 | 1.3 (3) | C32—C31—C36—C35 | 0.4 (3) |
| C12-C13-C14-F14 | 177.30 (15) | C43—C31—C36—C35 | -177.98 (18) |
| C12—C13—C14—C15 | -1.7 (3) | C56—N51—C52—C53 | 1.3 (3) |
| C13—C14—C15—C16 | 1.0 (3) | N51—C52—C53—C54 | -0.3 (3) |
| F14-C14-C15-C16 | -177.96 (16) | N51—C52—C53—C57 | 178.99 (18) |
| C12—C11—C16—C15 | -0.4 (3) | C52—C53—C54—C55 | -1.3 (3) |
| C1-C11-C16-C15 | -176.87 (15) | C57—C53—C54—C55 | 179.41 (17) |
| C14—C15—C16—C11 | 0.1 (3) | C53—C54—C55—C56 | 1.9 (3) |
| N1-C1-C21-C26 | -111.11 (17) | C52—N51—C56—C55 | -0.6 (3) |
| C11—C1—C21—C26 | 125.50 (17) | C54—C55—C56—N51 | -0.9 (3) |
| N1—C1—C21—C22 | 69.4 (2) | C54—C53—C57—O52 | 2.9 (2) |
| C11—C1—C21—C22 | -54.0 (2) | C52—C53—C57—O52 | -176.37 (17) |
| C26—C21—C22—C23 | -0.3 (3) | C54—C53—C57—O51 | -176.63 (16) |
| C1—C21—C22—C23 | 179.18 (16) | C52—C53—C57—O51 | 4.1 (2) |
| | | | |

Hydrogen-bond geometry (Å, °)

Cg1 and Cg2 represent the centroids of the C11-C16 and C31-C36 rings, respectively.

| D—H···A | <i>D</i> —Н | H···A | D··· A | D—H··· A |
|--------------------------------------|-------------|-------|-------------|------------|
| N4—H4…O51 | 0.98 | 1.58 | 2.5630 (19) | 175 |
| $C3$ — $H3B$ ···· $O52^{i}$ | 0.99 | 2.53 | 3.440 (2) | 153 |
| С13—Н13…О52 ^{іі} | 0.95 | 2.57 | 3.442 (2) | 152 |
| C56—H56…O51 ⁱⁱⁱ | 0.95 | 2.34 | 3.236 (3) | 158 |
| C26—H26…Cg1 ^{iv} | 0.95 | 2.81 | 3.758 (2) | 173 |
| C55—H55···· <i>Cg</i> 2 ^v | 0.95 | 2.85 | 3.567 (2) | 133 |
| | | | | |

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

$(II) \ 1-[Bis(4-fluorophenyl)methyl]-4-[(2E)-3-phenylprop-2-en-1-yl] piperazine-1, 4-diium \ bis(4-bis(4-fluorophenyl)methyl)methyl] \ -4-[(2E)-3-phenylprop-2-en-1-yl] \ piperazine-1, 4-diium \ bis(4-bis(4-fluorophenyl)methyl)methyl] \ -4-[(2E)-3-phenylprop-2-en-1-yl] \ piperazine-1, 4-diium \ bis(4-bis(4-fluorophenyl)methyl)methyl] \ -4-[(2E)-3-phenylprop-2-en-1-yl] \ piperazine-1, 4-diium \ bis(4-bis(4-fluorophenyl)methyl] \ -4-[(2E)-3-phenylprop-2-en-1-yl] \ piperazine-1, 4-diium \ bis(4-bis(4-fluorophenylprop-2-en-1-yl] \ piperazine-1, 4-diium \ bis(4-bis(4-fluoro$

methylbenzenesulfonate) dihydrate

| Crystal data | |
|---|---|
| $C_{26}H_{28}F_2N_2{}^{2+}\cdot 2C_7H_7O_3S^-\cdot 2H_2O$ | V = 3853.2 (3) Å ³ |
| $M_r = 784.91$ | Z = 4 |
| Monoclinic, $P2_1/c$ | F(000) = 1656 |
| a = 10.0546 (5) Å | $D_{\rm x} = 1.353 {\rm ~Mg} {\rm ~m}^{-3}$ |
| b = 14.8338 (6) Å | Mo <i>K</i> α radiation, $\lambda = 0.71073$ Å |
| c = 26.9437 (12) Å | Cell parameters from 9580 reflections |
| $\beta = 106.497 \ (3)^{\circ}$ | $\theta = 1.6 - 28.4^{\circ}$ |

 $\mu = 0.20 \text{ mm}^{-1}$ T = 200 K

Data collection

| Bruker APEXII CCD area-detector | 7212 independent reflections |
|--|---|
| diffractometer | 6219 reflections with $I > 2\sigma(I)$ |
| φ and ω scans | $R_{\rm int} = 0.038$ |
| Absorption correction: multi-scan | $\theta_{\rm max} = 25.6^{\circ}, \ \theta_{\rm min} = 2.1^{\circ}$ |
| (SADABS; Sheldrick, 2003) | $h = -12 \rightarrow 12$ |
| $T_{\min} = 0.877, \ T_{\max} = 0.953$ | $k = -15 \rightarrow 18$ |
| 45646 measured reflections | $l = -32 \rightarrow 32$ |
| | |
| Potinomont | |

Refinement

| Refinement on F^2 | Hydrogen site location: mixed |
|---------------------------------|--|
| Least-squares matrix: full | H-atom parameters constrained |
| $R[F^2 > 2\sigma(F^2)] = 0.075$ | $w = 1/[\sigma^2(F_o^2) + (0.0731P)^2 + 8.0533P]$ |
| $wR(F^2) = 0.202$ | where $P = (F_{o}^{2} + 2F_{c}^{2})/3$ |
| S = 1.09 | $(\Delta/\sigma)_{\rm max} = 0.001$ |
| 7212 reflections | $\Delta \rho_{\rm max} = 0.71 \text{ e } \text{\AA}^{-3}$ |
| 527 parameters | $\Delta \rho_{\rm min} = -0.60 \text{ e } \text{\AA}^{-3}$ |
| 27 restraints | |
| | |

Special details

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

Block, yellow

 $0.59 \times 0.43 \times 0.24 \text{ mm}$

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

| | x | у | Ζ | $U_{ m iso}$ */ $U_{ m eq}$ | Occ. (<1) |
|-----|------------|--------------|--------------|-----------------------------|-----------|
| N1 | 0.9062 (3) | 0.31590 (17) | 0.06845 (9) | 0.0283 (5) | |
| H1 | 0.9988 | 0.3286 | 0.0707 | 0.034* | |
| C2 | 0.8798 (4) | 0.3563 (2) | 0.11553 (12) | 0.0377 (8) | |
| H2A | 0.9473 | 0.3313 | 0.1467 | 0.045* | |
| H2B | 0.7857 | 0.3391 | 0.1167 | 0.045* | |
| C3 | 0.8915 (3) | 0.4569 (2) | 0.11658 (12) | 0.0329 (7) | |
| H3A | 0.8675 | 0.4802 | 0.1474 | 0.040* | |
| H3B | 0.9888 | 0.4740 | 0.1198 | 0.040* | |
| N4 | 0.7998 (3) | 0.49940 (19) | 0.06970 (10) | 0.0374 (6) | |
| H4 | 0.7026 | 0.4838 | 0.0691 | 0.045* | |
| C5 | 0.8239 (4) | 0.4592 (2) | 0.02266 (12) | 0.0376 (8) | |
| H5A | 0.9185 | 0.4751 | 0.0213 | 0.045* | |
| H5B | 0.7570 | 0.4853 | -0.0083 | 0.045* | |
| C6 | 0.8088 (4) | 0.3587 (2) | 0.02114 (13) | 0.0421 (8) | |
| H6A | 0.8289 | 0.3351 | -0.0103 | 0.050* | |
| H6B | 0.7120 | 0.3426 | 0.0193 | 0.050* | |
| C1 | 0.8877 (4) | 0.2133 (2) | 0.06603 (13) | 0.0376 (8) | |
| H1A | 0.7879 | 0.2008 | 0.0629 | 0.045* | |
| C11 | 0.9731 (4) | 0.1704 (2) | 0.11671 (14) | 0.0411 (8) | |
| C12 | 0.9029 (4) | 0.1445 (3) | 0.15121 (15) | 0.0464 (9) | |
| | | | | | |

| H12 | 0.8061 | 0.1555 | 0.1435 | 0.056* |
|--------------------------|------------|--------------|---------------|-----------------|
| C13 | 0.9716 (5) | 0.1025 (3) | 0.19714 (16) | 0.0549 (10) |
| H13 | 0.9228 | 0.0848 | 0.2210 | 0.066* |
| C14 | 1.1094 (5) | 0.0870 (3) | 0.20764 (16) | 0.0572 (11) |
| F14 | 1.1764 (3) | 0.04445 (19) | 0.25255 (11) | 0.0858 (10) |
| C15 | 1.1850 (5) | 0.1137 (3) | 0.17454 (19) | 0.0610(12) |
| H15 | 1.2822 | 0.1039 | 0.1830 | 0.073* |
| C16 | 1.1131 (4) | 0.1556 (3) | 0.12814 (16) | 0.0491 (9) |
| H16 | 1.1618 | 0.1740 | 0.1044 | 0.059* |
| C21 | 0.9177 (4) | 0.1744(2) | 0.01823 (13) | 0.0377 (8) |
| C22 | 1 0266 (4) | 0.1995(3) | -0.00073(16) | 0.0511(10) |
| H22 | 1.0200 (1) | 0.2450 | 0.0167 | 0.061* |
| C23 | 1.0050 | 0.1601 (3) | -0.04430(16) | 0.0542(10) |
| H23 | 1 1211 | 0.1780 | -0.0571 | 0.0542 (10) |
| C24 | 0.9575(5) | 0.0957 (3) | -0.06822(14) | 0.000 |
| E24 | 0.9373(3) | 0.05713(10) | -0.11165(10) | 0.0303(10) |
| 1 ⁻ 24 C25 | 0.9777(4) | 0.05713(19) | -0.05212(16) | 0.0848(10) |
| C25 | 0.8303 (3) | 0.0085 (5) | -0.03213(10) | 0.0388 (11) |
| H23 | 0.7903 | 0.0222 | -0.0701 | 0.071° |
| C26 | 0.8279 (4) | 0.1087 (5) | -0.008/2 (16) | 0.0504 (9) |
| H26 | 0.7504 | 0.0910 | 0.0026 | 0.061* |
| C41 | 0.8089 (4) | 0.5998 (2) | 0.06908 (14) | 0.0440 (8) |
| H4IA | 0.9029 | 0.6170 | 0.0677 | 0.053* |
| H41B | 0.7418 | 0.6225 | 0.0371 | 0.053* |
| C42 | 0.7812 (4) | 0.6449 (2) | 0.11408 (14) | 0.0432 (8) |
| H42 | 0.6939 | 0.6343 | 0.1201 | 0.052* |
| C43 | 0.8675 (4) | 0.6978 (3) | 0.14586 (15) | 0.0457 (9) |
| H43 | 0.9579 | 0.7021 | 0.1416 | 0.055* |
| C31 | 0.8407 (4) | 0.7523 (3) | 0.18798 (14) | 0.0442 (8) |
| C32 | 0.9371 (5) | 0.8177 (3) | 0.21029 (16) | 0.0580 (11) |
| H32 | 1.0175 | 0.8247 | 0.1988 | 0.070* |
| C33 | 0.9191 (6) | 0.8722 (3) | 0.24818 (18) | 0.0676 (13) |
| H33 | 0.9861 | 0.9174 | 0.2622 | 0.081* |
| C34 | 0.8081 (6) | 0.8634 (3) | 0.26634 (17) | 0.0710 (15) |
| H34 | 0.7970 | 0.9018 | 0.2931 | 0.085* |
| C35 | 0.7098 (5) | 0.7972 (4) | 0.24544 (19) | 0.0681 (13) |
| H35 | 0.6314 | 0.7895 | 0.2581 | 0.082* |
| C36 | 0.7275 (4) | 0.7427 (3) | 0.20607 (16) | 0.0527 (10) |
| H36 | 0.6600 | 0.6981 | 0.1914 | 0.063* |
| C51 | 1.3052 (3) | 0.6677 (2) | 0.17881 (13) | 0.0366 (7) |
| C52 | 1.2814 (4) | 0.7583 (3) | 0.16840 (15) | 0.0446 (8) |
| H52 | 1.2474 | 0.7782 | 0.1336 | 0.054* |
| C53 | 1 3065 (4) | 0.8197(3) | 0 20801 (16) | 0.0507 (9) |
| Н53 | 1.2896 | 0.8818 | 0.2003 | 0.061* |
| C54 | 1.3564 (4) | 0.7923 (3) | 0.25945 (16) | 0.0504 (9) |
| C55 | 1 3787 (5) | 0.7012(3) | 0 26940 (15) | 0.0523(10) |
| H55 | 1 4118 | 0.6810 | 0 3042 | 0.063* |
| C56 | 1 3537 (4) | 0.6388 (3) | 0.22950 (14) | 0.0484 (9) |
| US0 H56 | 1 3698 | 0.5765 | 0.22750 (14) | 0.058* |
| 11.0 | 1.5070 | 0.0700 | 0.2370 | 0.000 |

| C57 | 1.3887 (6) | 0.8599 (4) | 0.30303 (19) | 0.0741 (14) | |
|------|--------------|--------------|---------------|-------------|-----------|
| H57A | 1.3970 | 0.8287 | 0.3358 | 0.111* | |
| H57B | 1.3139 | 0.9045 | 0.2970 | 0.111* | |
| H57C | 1.4763 | 0.8904 | 0.3046 | 0.111* | |
| S51 | 1.27821 (10) | 0.59086 (6) | 0.12647 (3) | 0.0400 (2) | |
| O51 | 1.2760 (4) | 0.50202 (19) | 0.14856 (11) | 0.0621 (8) | |
| O52 | 1.3909 (3) | 0.60300 (19) | 0.10373 (12) | 0.0601 (8) | |
| O53 | 1.1460 (3) | 0.6137 (2) | 0.09039 (10) | 0.0662 (9) | |
| C61 | 0.4319 (4) | 0.1735 (3) | -0.00551 (18) | 0.0413 (11) | 0.832 (6) |
| C62 | 0.3808 (17) | 0.0869 (7) | -0.0143 (4) | 0.093 (2) | 0.832 (6) |
| H62 | 0.3444 | 0.0576 | 0.0104 | 0.112* | 0.832 (6) |
| C63 | 0.3830 (13) | 0.0428 (6) | -0.0596 (4) | 0.092 (4) | 0.832 (6) |
| H63 | 0.3522 | -0.0180 | -0.0645 | 0.110* | 0.832 (6) |
| C64 | 0.4279 (13) | 0.0839 (6) | -0.0970 (3) | 0.0534 (13) | 0.832 (6) |
| C65 | 0.4758 (6) | 0.1721 (4) | -0.0882(2) | 0.0484 (15) | 0.832 (6) |
| H65 | 0.5067 | 0.2027 | -0.1138 | 0.058* | 0.832 (6) |
| C66 | 0.4789 (5) | 0.2158 (3) | -0.04249 (19) | 0.0450 (13) | 0.832 (6) |
| H66 | 0.5139 | 0.2755 | -0.0367 | 0.054* | 0.832 (6) |
| C67 | 0.4262 (16) | 0.0347 (6) | -0.1462 (4) | 0.070(2) | 0.832 (6) |
| H67A | 0.4667 | 0.0732 | -0.1676 | 0.105* | 0.832 (6) |
| H67B | 0.3303 | 0.0199 | -0.1652 | 0.105* | 0.832 (6) |
| H67C | 0.4804 | -0.0209 | -0.1376 | 0.105* | 0.832 (6) |
| S61 | 0.44568 (17) | 0.22444 (13) | 0.05538 (6) | 0.0442 (4) | 0.832 (6) |
| O61 | 0.3748 (4) | 0.3104 (2) | 0.04318 (14) | 0.0566 (11) | 0.832 (6) |
| O62 | 0.3801 (4) | 0.1655 (3) | 0.08433 (15) | 0.0681 (12) | 0.832 (6) |
| O63 | 0.5873 (5) | 0.2398 (5) | 0.0767 (2) | 0.097 (2) | 0.832 (6) |
| C61A | 0.474 (2) | 0.1737 (10) | -0.0054 (6) | 0.0413 (11) | 0.168 (6) |
| C62A | 0.397 (8) | 0.095 (3) | -0.0121 (15) | 0.093 (2) | 0.168 (6) |
| H62A | 0.3742 | 0.0678 | 0.0164 | 0.112* | 0.168 (6) |
| C63A | 0.353 (7) | 0.056 (4) | -0.0611 (16) | 0.092 (4) | 0.168 (6) |
| H63A | 0.2792 | 0.0139 | -0.0684 | 0.110* | 0.168 (6) |
| C64A | 0.413 (7) | 0.078 (3) | -0.0988 (13) | 0.0534 (13) | 0.168 (6) |
| C65A | 0.502 (4) | 0.152 (2) | -0.0898 (8) | 0.0484 (15) | 0.168 (6) |
| H65A | 0.5517 | 0.1670 | -0.1138 | 0.058* | 0.168 (6) |
| C66A | 0.518 (3) | 0.2049 (15) | -0.0460 (7) | 0.0450 (13) | 0.168 (6) |
| H66A | 0.5591 | 0.2628 | -0.0441 | 0.054* | 0.168 (6) |
| C67A | 0.405 (9) | 0.015 (4) | -0.1434 (19) | 0.070 (2) | 0.168 (6) |
| H67D | 0.3240 | 0.0304 | -0.1723 | 0.105* | 0.168 (6) |
| H67E | 0.3960 | -0.0471 | -0.1324 | 0.105* | 0.168 (6) |
| H67F | 0.4894 | 0.0206 | -0.1544 | 0.105* | 0.168 (6) |
| S61A | 0.4738 (9) | 0.2443 (6) | 0.0478 (3) | 0.0442 (4) | 0.168 (6) |
| O61A | 0.4807 (16) | 0.3360 (7) | 0.0291 (6) | 0.059 (5)* | 0.168 (6) |
| O62A | 0.3450 (13) | 0.2298 (11) | 0.0612 (6) | 0.067 (6)* | 0.168 (6) |
| O63A | 0.5974 (10) | 0.2273 (9) | 0.0855 (5) | 0.017 (3)* | 0.168 (6) |
| O71 | 1.1663 (3) | 0.37595 (19) | 0.07577 (12) | 0.0563 (7) | (-) |
| H71A | 1.1873 | 0.4171 | 0.0985 | 0.084* | |
| H71B | 1.2430 | 0.3652 | 0.0700 | 0.090* | |
| O81 | 0.5371 (3) | 0.4699 (2) | 0.07130 (16) | 0.0782 (11) | |

supporting information

| H81B | 0.4924 | 0.5051 | 0.0850 | 0.117* |
|------|--------|--------|--------|--------|
| H81A | 0.4824 | 0.4292 | 0.0563 | 0.117* |

Atomic displacement parameters $(Å^2)$

| | U^{11} | U^{22} | U^{33} | U^{12} | U^{13} | U^{23} |
|-----|-------------|-------------|-------------|--------------|--------------|--------------|
| N1 | 0.0293 (13) | 0.0278 (13) | 0.0288 (13) | -0.0005 (10) | 0.0099 (10) | -0.0034 (10) |
| C2 | 0.050(2) | 0.0381 (18) | 0.0283 (16) | 0.0069 (15) | 0.0164 (15) | -0.0004 (13) |
| C3 | 0.0369 (17) | 0.0332 (17) | 0.0273 (15) | 0.0053 (13) | 0.0070 (13) | -0.0024 (13) |
| N4 | 0.0484 (17) | 0.0359 (15) | 0.0300 (14) | 0.0117 (13) | 0.0144 (12) | 0.0002 (12) |
| C5 | 0.0432 (19) | 0.0415 (19) | 0.0284 (16) | 0.0103 (15) | 0.0108 (14) | -0.0007 (14) |
| C6 | 0.049 (2) | 0.044 (2) | 0.0312 (17) | 0.0087 (16) | 0.0092 (15) | -0.0010 (15) |
| C1 | 0.0378 (18) | 0.0334 (18) | 0.0418 (19) | -0.0049 (14) | 0.0114 (15) | -0.0050 (14) |
| C11 | 0.053 (2) | 0.0281 (17) | 0.0437 (19) | 0.0048 (15) | 0.0162 (17) | -0.0046 (14) |
| C12 | 0.056 (2) | 0.039 (2) | 0.047 (2) | 0.0028 (17) | 0.0188 (18) | 0.0022 (16) |
| C13 | 0.076 (3) | 0.042 (2) | 0.049 (2) | 0.002 (2) | 0.022 (2) | 0.0085 (18) |
| C14 | 0.082 (3) | 0.033 (2) | 0.045 (2) | 0.003 (2) | 0.001 (2) | 0.0006 (17) |
| F14 | 0.110 (2) | 0.0607 (17) | 0.0622 (16) | 0.0123 (16) | -0.0149 (16) | 0.0141 (13) |
| C15 | 0.046 (2) | 0.045 (2) | 0.083 (3) | 0.0035 (18) | 0.004 (2) | -0.012 (2) |
| C16 | 0.056 (2) | 0.037 (2) | 0.056 (2) | -0.0032 (17) | 0.0189 (19) | -0.0028 (17) |
| C21 | 0.049 (2) | 0.0317 (17) | 0.0335 (17) | 0.0078 (15) | 0.0142 (15) | -0.0001 (14) |
| C22 | 0.056 (2) | 0.047 (2) | 0.055 (2) | -0.0161 (18) | 0.0226 (19) | -0.0180 (18) |
| C23 | 0.074 (3) | 0.049 (2) | 0.052 (2) | -0.002 (2) | 0.037 (2) | -0.0044 (18) |
| C24 | 0.079 (3) | 0.037 (2) | 0.0364 (19) | 0.0094 (19) | 0.0184 (19) | -0.0044 (15) |
| F24 | 0.150 (3) | 0.0631 (17) | 0.0510 (15) | -0.0019 (18) | 0.0443 (17) | -0.0222 (13) |
| C25 | 0.075 (3) | 0.044 (2) | 0.052 (2) | -0.015 (2) | 0.009 (2) | -0.0171 (19) |
| C26 | 0.053 (2) | 0.040 (2) | 0.061 (2) | -0.0046 (17) | 0.022 (2) | 0.0034 (18) |
| C41 | 0.056 (2) | 0.0346 (18) | 0.0409 (19) | 0.0125 (16) | 0.0140 (17) | 0.0048 (15) |
| C42 | 0.051 (2) | 0.0348 (19) | 0.045 (2) | 0.0131 (16) | 0.0159 (17) | 0.0001 (15) |
| C43 | 0.044 (2) | 0.045 (2) | 0.051 (2) | 0.0082 (17) | 0.0189 (17) | 0.0088 (17) |
| C31 | 0.053 (2) | 0.042 (2) | 0.0365 (18) | 0.0099 (17) | 0.0110 (16) | 0.0059 (15) |
| C32 | 0.063 (3) | 0.062 (3) | 0.050 (2) | -0.002 (2) | 0.016 (2) | 0.010 (2) |
| C33 | 0.090 (4) | 0.050 (3) | 0.051 (2) | -0.011 (2) | 0.001 (2) | 0.000 (2) |
| C34 | 0.112 (4) | 0.056 (3) | 0.040 (2) | 0.031 (3) | 0.013 (3) | -0.006 (2) |
| C35 | 0.064 (3) | 0.083 (3) | 0.065 (3) | 0.027 (3) | 0.032 (2) | 0.016 (3) |
| C36 | 0.053 (2) | 0.048 (2) | 0.054 (2) | 0.0001 (18) | 0.0090 (19) | 0.0028 (18) |
| C51 | 0.0362 (17) | 0.0393 (18) | 0.0359 (17) | 0.0018 (14) | 0.0130 (14) | 0.0037 (14) |
| C52 | 0.052 (2) | 0.041 (2) | 0.0402 (19) | 0.0053 (16) | 0.0121 (16) | 0.0039 (16) |
| C53 | 0.056 (2) | 0.040 (2) | 0.059 (2) | 0.0058 (17) | 0.022 (2) | 0.0019 (18) |
| C54 | 0.054 (2) | 0.054 (2) | 0.049 (2) | -0.0008 (18) | 0.0243 (18) | -0.0107 (18) |
| C55 | 0.065 (3) | 0.059 (2) | 0.0348 (19) | -0.005 (2) | 0.0172 (18) | 0.0042 (17) |
| C56 | 0.064 (2) | 0.044 (2) | 0.0393 (19) | -0.0007 (18) | 0.0185 (18) | 0.0058 (16) |
| C57 | 0.095 (4) | 0.071 (3) | 0.061 (3) | -0.003 (3) | 0.030 (3) | -0.019 (2) |
| S51 | 0.0485 (5) | 0.0403 (5) | 0.0339 (4) | -0.0020 (4) | 0.0159 (4) | 0.0013 (3) |
| O51 | 0.100 (2) | 0.0417 (15) | 0.0421 (15) | -0.0197 (15) | 0.0159 (15) | -0.0003 (12) |
| O52 | 0.0704 (19) | 0.0503 (17) | 0.076 (2) | -0.0065 (14) | 0.0471 (17) | -0.0141 (14) |
| O53 | 0.0642 (19) | 0.093 (2) | 0.0365 (14) | 0.0181 (17) | 0.0066 (13) | -0.0071 (15) |
| C61 | 0.021 (3) | 0.045 (2) | 0.060(2) | 0.0073 (18) | 0.0144 (19) | -0.0026 (18) |

| C62 | 0.153 (7) | 0.058 (4) | 0.100 (4) | -0.043 (4) | 0.088 (5) | -0.025 (3) |
|------|-------------|-------------|-------------|--------------|-------------|--------------|
| C63 | 0.160 (9) | 0.046 (4) | 0.094 (4) | -0.028 (6) | 0.076 (5) | -0.019 (3) |
| C64 | 0.057 (4) | 0.050 (3) | 0.053 (2) | 0.013 (2) | 0.014 (2) | 0.0022 (19) |
| C65 | 0.040 (3) | 0.057 (4) | 0.044 (2) | 0.006 (3) | 0.0053 (19) | 0.011 (2) |
| C66 | 0.030 (3) | 0.044 (2) | 0.054 (2) | 0.001 (2) | -0.001 (2) | 0.0033 (19) |
| C67 | 0.093 (7) | 0.062 (6) | 0.054 (3) | 0.010 (4) | 0.018 (3) | 0.000 (4) |
| S61 | 0.0298 (7) | 0.0456 (8) | 0.0606 (7) | -0.0028 (6) | 0.0183 (5) | -0.0034 (6) |
| O61 | 0.058 (2) | 0.053 (2) | 0.060 (2) | 0.0118 (16) | 0.0186 (17) | -0.0062 (16) |
| O62 | 0.072 (3) | 0.073 (3) | 0.068 (2) | 0.000 (2) | 0.034 (2) | 0.008 (2) |
| O63 | 0.068 (3) | 0.138 (6) | 0.091 (4) | 0.017 (3) | 0.034 (3) | -0.017 (4) |
| C61A | 0.021 (3) | 0.045 (2) | 0.060 (2) | 0.0073 (18) | 0.0144 (19) | -0.0026 (18) |
| C62A | 0.153 (7) | 0.058 (4) | 0.100 (4) | -0.043 (4) | 0.088 (5) | -0.025 (3) |
| C63A | 0.160 (9) | 0.046 (4) | 0.094 (4) | -0.028 (6) | 0.076 (5) | -0.019 (3) |
| C64A | 0.057 (4) | 0.050 (3) | 0.053 (2) | 0.013 (2) | 0.014 (2) | 0.0022 (19) |
| C65A | 0.040 (3) | 0.057 (4) | 0.044 (2) | 0.006 (3) | 0.0053 (19) | 0.011 (2) |
| C66A | 0.030 (3) | 0.044 (2) | 0.054 (2) | 0.001 (2) | -0.001 (2) | 0.0033 (19) |
| C67A | 0.093 (7) | 0.062 (6) | 0.054 (3) | 0.010 (4) | 0.018 (3) | 0.000 (4) |
| S61A | 0.0298 (7) | 0.0456 (8) | 0.0606 (7) | -0.0028 (6) | 0.0183 (5) | -0.0034 (6) |
| O71 | 0.0462 (15) | 0.0511 (16) | 0.0750 (19) | -0.0082 (12) | 0.0228 (14) | -0.0225 (14) |
| O81 | 0.0590 (19) | 0.065 (2) | 0.128 (3) | 0.0021 (16) | 0.056 (2) | -0.015 (2) |
| | | | | | | |

Geometric parameters (Å, °)

| N1—C2 | 1.493 (4) | С36—Н36 | 0.9500 |
|---------|-----------|----------|-----------|
| N1—C6 | 1.509 (4) | C51—C52 | 1.380 (5) |
| N1-C1 | 1.532 (4) | C51—C56 | 1.382 (5) |
| N1—H1 | 0.9355 | C51—S51 | 1.773 (4) |
| C2—C3 | 1.497 (5) | C52—C53 | 1.371 (5) |
| C2—H2A | 0.9900 | С52—Н52 | 0.9500 |
| C2—H2B | 0.9900 | C53—C54 | 1.394 (6) |
| C3—N4 | 1.477 (4) | С53—Н53 | 0.9500 |
| С3—НЗА | 0.9900 | C54—C55 | 1.384 (6) |
| С3—Н3В | 0.9900 | C54—C57 | 1.507 (6) |
| N4—C5 | 1.482 (4) | C55—C56 | 1.387 (6) |
| N4—C41 | 1.493 (4) | С55—Н55 | 0.9500 |
| N4—H4 | 0.9998 | С56—Н56 | 0.9500 |
| С5—С6 | 1.497 (5) | C57—H57A | 0.9800 |
| С5—Н5А | 0.9900 | С57—Н57В | 0.9800 |
| С5—Н5В | 0.9900 | С57—Н57С | 0.9800 |
| С6—Н6А | 0.9900 | S51—O52 | 1.444 (3) |
| С6—Н6В | 0.9900 | S51—O53 | 1.446 (3) |
| C1—C21 | 1.518 (5) | S51—O51 | 1.449 (3) |
| C1—C11 | 1.531 (5) | C61—C66 | 1.370 (6) |
| C1—H1A | 1.0000 | C61—C62 | 1.379 (7) |
| C11—C16 | 1.370 (5) | C61—S61 | 1.776 (4) |
| C11—C12 | 1.372 (5) | C62—C63 | 1.390 (8) |
| C12—C13 | 1.382 (6) | С62—Н62 | 0.9500 |
| C12—H12 | 0.9500 | C63—C64 | 1.361 (7) |
| | | | |

| C13—C14 | 1.352 (7) | С63—Н63 | 0.9500 |
|---------------------|-----------|----------------------------|---------------------|
| С13—Н13 | 0.9500 | C64—C65 | 1.390 (7) |
| C14—F14 | 1.361 (5) | C64—C67 | 1.508 (6) |
| C14—C15 | 1.384 (7) | C65—C66 | 1.384 (7) |
| C15—C16 | 1.400 (6) | С65—Н65 | 0.9500 |
| С15—Н15 | 0.9500 | С66—Н66 | 0.9500 |
| С16—Н16 | 0.9500 | С67—Н67А | 0.9800 |
| C21—C22 | 1.384 (5) | С67—Н67В | 0.9800 |
| C21—C26 | 1.386 (5) | С67—Н67С | 0.9800 |
| C22—C23 | 1.376 (5) | S61—O63 | 1.395 (5) |
| С22—Н22 | 0.9500 | S61—O62 | 1.448 (4) |
| C23—C24 | 1.341 (6) | S61—O61 | 1.452 (4) |
| С23—Н23 | 0.9500 | C61A—C66A | 1.373 (7) |
| C24—C25 | 1.332 (6) | C61A—C62A | 1.383 (8) |
| C24—F24 | 1.368 (4) | C61A—S61A | 1.773 (6) |
| C_{25} C_{26} | 1.388 (6) | C62A—C63A | 1.392 (10) |
| C25—H25 | 0.9500 | C62A - H62A | 0.9500 |
| C26—H26 | 0.9500 | C63A—C64A | 1.362 (8) |
| C41-C42 | 1,479 (5) | C63A—H63A | 0.9500 |
| C41—H41A | 0.9900 | C64A—C65A | 1.392 (9) |
| C41—H41B | 0.9900 | C64A—C67A | 1.509 (8) |
| C42—C43 | 1,297 (6) | C65A—C66A | 1.384 (8) |
| C42—H42 | 0.9500 | C65A—H65A | 0.9500 |
| C43-C31 | 1,479 (5) | C66A—H66A | 0.9500 |
| C43—H43 | 0.9500 | C67A—H67D | 0.9800 |
| C31—C36 | 1.367 (6) | С67А—Н67Е | 0.9800 |
| C31—C32 | 1.382 (6) | C67A—H67F | 0.9800 |
| C32—C33 | 1.353 (7) | S61A-063A | 1.388 (6) |
| С32—Н32 | 0.9500 | S61A—062A | 1.456 (6) |
| C33—C34 | 1.346 (8) | S61A—O61A | 1.458 (6) |
| С33—Н33 | 0.9500 | 071—H71A | 0.8475 |
| C34—C35 | 1.394 (8) | 071—H71B | 0.8448 |
| С34—Н34 | 0.9500 | O81—H81B | 0.8387 |
| C35—C36 | 1.384 (6) | 081—H81A | 0.8393 |
| С35—Н35 | 0.9500 | | |
| | | | |
| C2—N1—C6 | 108.6 (2) | С33—С34—Н34 | 120.4 |
| C_2 N1-C1 | 112.7 (2) | C35—C34—H34 | 120.4 |
| C6-N1-C1 | 109.9(2) | $C_{36} - C_{35} - C_{34}$ | 1193(4) |
| C2—N1—H1 | 106.2 | C36—C35—H35 | 120.3 |
| C6—N1—H1 | 111.4 | C34—C35—H35 | 120.3 |
| C1—N1—H1 | 108.0 | $C_{31} - C_{36} - C_{35}$ | 120.9 (4) |
| N1-C2-C3 | 112.6 (3) | C31—C36—H36 | 119.5 |
| N1-C2-H2A | 109.1 | C35—C36—H36 | 119.5 |
| C3 - C2 - H2A | 109.1 | C52—C51—C56 | 119.7 (3) |
| N1—C2—H2B | 109.1 | C52—C51—S51 | 119.1 (3) |
| C3 - C2 - H2B | 109.1 | C56—C51—S51 | 121.2(3) |
| $H^2A - C^2 - H^2B$ | 107.8 | C_{53} C_{52} C_{51} | 121.2(3) 1204(3) |
| | 107.0 | 000 002 001 | 120.1 (3) |

| N4—C3—C2 | 112.5 (3) | С53—С52—Н52 | 119.8 |
|-------------------------------------|----------------------|----------------------------|--------------------------|
| N4—C3—H3A | 109.1 | C51—C52—H52 | 119.8 |
| С2—С3—НЗА | 109.1 | C52—C53—C54 | 121.1 (4) |
| N4—C3—H3B | 109.1 | С52—С53—Н53 | 119.5 |
| С2—С3—Н3В | 109.1 | С54—С53—Н53 | 119.5 |
| НЗА—СЗ—НЗВ | 107.8 | C55—C54—C53 | 118.0 (4) |
| C3—N4—C5 | 110.2 (2) | C55—C54—C57 | 120.7 (4) |
| C3—N4—C41 | 114.2 (3) | C53—C54—C57 | 121.2 (4) |
| C5—N4—C41 | 111.5 (3) | C54—C55—C56 | 121.2 (4) |
| C3—N4—H4 | 106.5 | C54—C55—H55 | 119.4 |
| C5—N4—H4 | 106.9 | С56—С55—Н55 | 119.4 |
| C41—N4—H4 | 107.0 | C51—C56—C55 | 119.6 (4) |
| N4 | 112.6 (3) | C51—C56—H56 | 120.2 |
| N4-C5-H5A | 109.1 | C55-C56-H56 | 120.2 |
| C6 | 109.1 | C54—C57—H57A | 109.5 |
| N4—C5—H5B | 109.1 | C54—C57—H57B | 109.5 |
| C6-C5-H5B | 109.1 | H57A_C57_H57B | 109.5 |
| H_{5A} C_{5} H_{5B} | 107.8 | C54—C57—H57C | 109.5 |
| C5-C6-N1 | 111 1 (3) | H57A_C57_H57C | 109.5 |
| C_{5} C_{6} H_{6} | 109.4 | H57B_C57_H57C | 109.5 |
| N1-C6-H6A | 109.4 | 052 - 851 - 053 | 111 57 (19) |
| C5-C6-H6B | 109.4 | 052 551 055 | 111.37(19) 113.15(19) |
| N1-C6-H6B | 109.4 | 052 551 051 | 113.13(1) |
| | 109.4 | 053 551 051 | 107.21(17) |
| $C_{21} - C_{11} - C_{11}$ | 113.7(3) | 052 - 551 - 051 | 107.21(17) 106.72(17) |
| C_{21} C_{1} N_{1} | 113.7(3) | 055 551 051 | 105.94(16) |
| C_{11} C_{1} N_{1} | 111.3(3) 110.2(3) | $C_{66} - C_{61} - C_{62}$ | 119 4 (4) |
| C_{21} C_{1} H_{1A} | 107.1 | $C_{66} = C_{61} = S_{61}$ | 119.4(4) 121.7(3) |
| C_{11} C_{1} H_{1A} | 107.1 | C_{62} C_{61} S_{61} | 121.7(5) 118 8 (4) |
| N1 - C1 - H1A | 107.1 | C61 - C62 - C63 | 110.0(1) 119.4(5) |
| C_{16} | 119.8 (4) | C61 - C62 - H62 | 120.3 |
| C_{16} C_{11} C_{1} | 123.2(3) | C63 - C62 - H62 | 120.3 |
| C_{12} C_{11} C_{11} C_{11} | 123.2(3) 117.0(3) | C64 - C63 - C62 | 120.5 122.0(5) |
| C_{11} C_{12} C_{13} | 117.0(3) 120.7(4) | C64 - C63 - H63 | 119.0 |
| $C_{11} - C_{12} - H_{12}$ | 120.7 (4) | C62 - C63 - H63 | 119.0 |
| C_{13} C_{12} H_{12} | 119.7 | C63 - C64 - C65 | 117.9 (4) |
| C_{14} C_{13} C_{12} C_{12} | 119.1 (4) | C63 - C64 - C67 | 120.7(5) |
| $C_{14} - C_{13} - H_{13}$ | 120.5 | C65 - C64 - C67 | 120.7(5) 121.5(5) |
| C12_C13_H13 | 120.5 | C65 - C64 - C64 | 121.3(3) |
| C12 - C13 - H13 | 120.3 118 9 (4) | C66-C65-H65 | 119.6 |
| C_{13} C_{14} C_{15} | 110.9(4) 122.3(4) | C64—C65—H65 | 119.6 |
| F_{14} C_{14} C_{15} | 122.3(4) 118.8(4) | C_{61} C_{66} C_{65} | 120.5(4) |
| $C_{14} - C_{15} - C_{16}$ | 117.6(4) | C61 - C66 - H66 | 119 7 |
| C14-C15-H15 | 121.2 | C65—C66—H66 | 110.7 |
| C16—C15—H15 | 121.2 | $C64-C67-H67\Delta$ | 109 5 |
| C_{11} C_{16} C_{15} | 121.2 | C64—C67—H67B | 109.5 |
| C11—C16—H16 | 119 7 | H67A—C67—H67B | 109.5 |
| C15—C16—H16 | 119.7 | C64-C67-H67C | 109.5 |
| | · · / · / | | |

| C22—C21—C26 | 117.5 (3) | Н67А—С67—Н67С | 109.5 |
|---------------|------------|-----------------|------------|
| C22—C21—C1 | 125.4 (3) | Н67В—С67—Н67С | 109.5 |
| C26—C21—C1 | 117.1 (3) | O63—S61—O62 | 116.4 (4) |
| C23—C22—C21 | 121.5 (4) | O63—S61—O61 | 109.1 (4) |
| С23—С22—Н22 | 119.3 | O62—S61—O61 | 112.5 (2) |
| С21—С22—Н22 | 119.3 | O63—S61—C61 | 104.6 (3) |
| C24—C23—C22 | 118.3 (4) | O62—S61—C61 | 108.2 (2) |
| С24—С23—Н23 | 120.8 | O61—S61—C61 | 105.1 (2) |
| С22—С23—Н23 | 120.8 | C66A—C61A—C62A | 118.3 (7) |
| C25—C24—C23 | 123.3 (4) | C66A—C61A—S61A | 121.2 (7) |
| C25—C24—F24 | 118.8 (4) | C62A—C61A—S61A | 118.5 (8) |
| C23—C24—F24 | 117.9 (4) | C61A—C62A—C63A | 119.1 (9) |
| C24—C25—C26 | 119.0 (4) | C61A—C62A—H62A | 120.5 |
| C24—C25—H25 | 120.5 | C63A—C62A—H62A | 120.5 |
| C26—C25—H25 | 120.5 | C64A—C63A—C62A | 121.5 (11) |
| C21—C26—C25 | 120.3 (4) | С64А—С63А—Н63А | 119.3 |
| C21—C26—H26 | 119.8 | С62А—С63А—Н63А | 119.3 |
| С25—С26—Н26 | 119.8 | C63A—C64A—C65A | 117.1 (7) |
| C42—C41—N4 | 114.5 (3) | C63A—C64A—C67A | 120.7 (9) |
| C42—C41—H41A | 108.6 | C65A—C64A—C67A | 121.2 (11) |
| N4—C41—H41A | 108.6 | C66A—C65A—C64A | 120.4 (8) |
| C42—C41—H41B | 108.6 | С66А—С65А—Н65А | 119.8 |
| N4—C41—H41B | 108.6 | С64А—С65А—Н65А | 119.8 |
| H41A—C41—H41B | 107.6 | C61A—C66A—C65A | 120.4 (7) |
| C43—C42—C41 | 124.6 (4) | C61A—C66A—H66A | 119.8 |
| C43—C42—H42 | 117.7 | C65A—C66A—H66A | 119.8 |
| C41—C42—H42 | 117.7 | C64A—C67A—H67D | 109.5 |
| C42—C43—C31 | 127.2 (4) | С64А—С67А—Н67Е | 109.5 |
| С42—С43—Н43 | 116.4 | Н67D—С67А—Н67Е | 109.5 |
| C31—C43—H43 | 116.4 | C64A—C67A—H67F | 109.5 |
| C36—C31—C32 | 118.0 (4) | H67D—C67A—H67F | 109.5 |
| C36—C31—C43 | 124.6 (4) | H67E—C67A—H67F | 109.5 |
| C32—C31—C43 | 117.4 (4) | O63A—S61A—O62A | 117.7 (7) |
| C33—C32—C31 | 121.3 (4) | O63A—S61A—O61A | 107.6 (6) |
| С33—С32—Н32 | 119.3 | O62A—S61A—O61A | 110.9 (6) |
| С31—С32—Н32 | 119.3 | O63A—S61A—C61A | 106.3 (6) |
| C34—C33—C32 | 121.2 (5) | O62A—S61A—C61A | 108.4 (6) |
| С34—С33—Н33 | 119.4 | O61A—S61A—C61A | 105.2 (6) |
| С32—С33—Н33 | 119.4 | H71A—O71—H71B | 102.4 |
| C33—C34—C35 | 119.2 (4) | H81B—O81—H81A | 107.5 |
| | | | |
| C6—N1—C2—C3 | -55.3 (4) | C33—C34—C35—C36 | -0.8 (7) |
| C1—N1—C2—C3 | -177.3 (3) | C32—C31—C36—C35 | 0.2 (6) |
| N1-C2-C3-N4 | 55.4 (4) | C43—C31—C36—C35 | -179.4 (4) |
| C2—C3—N4—C5 | -53.2 (4) | C34—C35—C36—C31 | 0.8 (7) |
| C2—C3—N4—C41 | -179.7 (3) | C56—C51—C52—C53 | 0.4 (6) |
| C3—N4—C5—C6 | 54.7 (4) | S51—C51—C52—C53 | -177.5 (3) |
| C41—N4—C5—C6 | -177.3 (3) | C51—C52—C53—C54 | 0.0 (6) |
| | | | |

| | | | / |
|-----------------|------------|---------------------|-------------|
| N4—C5—C6—N1 | -57.2 (4) | C52—C53—C54—C55 | -0.6 (6) |
| C2—N1—C6—C5 | 55.9 (4) | C52—C53—C54—C57 | 177.9 (4) |
| C1—N1—C6—C5 | 179.6 (3) | C53—C54—C55—C56 | 0.6 (6) |
| C6—N1—C1—C21 | 59.7 (3) | C57—C54—C55—C56 | -177.8 (4) |
| C2—N1—C1—C11 | -51.9 (4) | C52—C51—C56—C55 | -0.4 (6) |
| C2—N1—C1—C21 | -179.0 (3) | S51—C51—C56—C55 | 177.5 (3) |
| C6—N1—C1—C11 | -173.2 (3) | C54—C55—C56—C51 | -0.2 (6) |
| C21—C1—C11—C16 | 46.3 (5) | C52—C51—S51—O52 | 73.3 (3) |
| N1-C1-C11-C16 | -79.4 (4) | C56—C51—S51—O52 | -104.6 (3) |
| C21—C1—C11—C12 | -132.4 (3) | C52—C51—S51—O53 | -46.4 (3) |
| N1-C1-C11-C12 | 101.8 (4) | C56-C51-S51-O53 | 135.8 (3) |
| C16—C11—C12—C13 | -0.9 (6) | C52—C51—S51—O51 | -165.6 (3) |
| C1—C11—C12—C13 | 177.9 (3) | C56—C51—S51—O51 | 16.5 (4) |
| C11—C12—C13—C14 | -0.2 (6) | C66—C61—C62—C63 | 2.5 (17) |
| C12-C13-C14-F14 | -179.0 (4) | S61—C61—C62—C63 | -173.3 (9) |
| C12—C13—C14—C15 | 1.6 (6) | C61—C62—C63—C64 | -3.3 (18) |
| C13—C14—C15—C16 | -2.0 (6) | C62—C63—C64—C65 | 1.7 (15) |
| F14—C14—C15—C16 | 178.7 (4) | C62—C63—C64—C67 | -178.7 (13) |
| C12—C11—C16—C15 | 0.5 (6) | C63—C64—C65—C66 | 0.7 (13) |
| C1-C11-C16-C15 | -178.2 (3) | C67—C64—C65—C66 | -178.9 (9) |
| C14—C15—C16—C11 | 0.8 (6) | C62—C61—C66—C65 | -0.2 (11) |
| C11—C1—C21—C22 | -82.7 (4) | S61—C61—C66—C65 | 175.5 (4) |
| N1-C1-C21-C22 | 42.4 (5) | C64—C65—C66—C61 | -1.4 (9) |
| C11—C1—C21—C26 | 97.7 (4) | C66—C61—S61—O63 | -58.5 (5) |
| N1-C1-C21-C26 | -137.2 (3) | C62—C61—S61—O63 | 117.3 (10) |
| C26—C21—C22—C23 | -0.8 (6) | C66—C61—S61—O62 | 176.8 (4) |
| C1—C21—C22—C23 | 179.6 (4) | C62—C61—S61—O62 | -7.5 (10) |
| C21—C22—C23—C24 | -0.4 (7) | C66—C61—S61—O61 | 56.4 (4) |
| C22—C23—C24—C25 | 0.5 (7) | C62—C61—S61—O61 | -127.8 (9) |
| C22—C23—C24—F24 | 179.7 (4) | C66A—C61A—C62A—C63A | -8 (7) |
| C23—C24—C25—C26 | 0.5 (7) | S61A—C61A—C62A—C63A | 157 (4) |
| F24—C24—C25—C26 | -178.7 (4) | C61A—C62A—C63A—C64A | 18 (7) |
| C22—C21—C26—C25 | 1.9 (6) | C62A—C63A—C64A—C65A | -12 (6) |
| C1—C21—C26—C25 | -178.5 (4) | C62A—C63A—C64A—C67A | 157 (7) |
| C24—C25—C26—C21 | -1.7 (7) | C63A—C64A—C65A—C66A | -5 (7) |
| C3—N4—C41—C42 | -56.2 (4) | C67A—C64A—C65A—C66A | -174 (5) |
| C5—N4—C41—C42 | 178.0 (3) | C62A—C61A—C66A—C65A | -9 (5) |
| N4—C41—C42—C43 | 121.6 (4) | S61A—C61A—C66A—C65A | -173 (3) |
| C41—C42—C43—C31 | 172.9 (3) | C64A—C65A—C66A—C61A | 16 (6) |
| C42—C43—C31—C36 | 12.6 (6) | C66A—C61A—S61A—O63A | -95 (2) |
| C42—C43—C31—C32 | -167.1 (4) | C62A—C61A—S61A—O63A | 101 (4) |
| C36—C31—C32—C33 | -1.3 (6) | C66A—C61A—S61A—O62A | 137 (2) |
| C43—C31—C32—C33 | 178.3 (4) | C62A—C61A—S61A—O62A | -27 (4) |
| C31—C32—C33—C34 | 1.4 (7) | C66A—C61A—S61A—O61A | 19 (2) |
| C32—C33—C34—C35 | -0.3 (7) | C62A—C61A—S61A—O61A | -145 (4) |
| | | | ~ / |

| D—H···A | <i>D</i> —Н | $H \cdots A$ | $D \cdots A$ | D—H···A |
|---------------------------------------|-------------|--------------|--------------|---------|
| N1—H1…O71 | 0.93 | 1.79 | 2.716 (4) | 169 |
| N4—H4…O81 | 1.00 | 1.69 | 2.690 (4) | 173 |
| O71—H71A···O51 | 0.85 | 1.88 | 2.708 (4) | 167 |
| O71—H71 <i>B</i> ···O61 ⁱ | 0.84 | 1.87 | 2.675 (5) | 160 |
| O81—H81A···O61 | 0.84 | 2.05 | 2.852 (5) | 161 |
| O81—H81 <i>B</i> ···O52 ⁱⁱ | 0.84 | 1.92 | 2.748 (4) | 167 |
| C1—H1A···O63 | 1.00 | 2.23 | 3.139 (7) | 151 |
| C2—H2 <i>B</i> ···O63 | 0.99 | 2.47 | 3.314 (7) | 143 |
| С6—Н6А…О53ііі | 0.99 | 2.37 | 3.191 (5) | 140 |
| C16—H16…O62 ⁱ | 0.95 | 2.41 | 3.228 (6) | 144 |
| C22—H22…O71 | 0.95 | 2.49 | 3.381 (5) | 156 |
| C34—H34…O51 ^{iv} | 0.95 | 2.42 | 3.362 (6) | 169 |
| C41—H41A····O53 | 0.99 | 2.35 | 3.280 (5) | 157 |
| C41—H41 <i>B</i> ···O61 ^v | 0.99 | 2.37 | 3.341 (5) | 167 |

Hydrogen-bond geometry (Å, °)

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