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1,3-Bis{(*E*)-[4-(dimethylamino)benzylidene]amino}propan-2-ol: chain structure formation *via* an O—H···N hydrogen bond

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The asymmetric unit of the title compound, $C_{21}H_{28}N_4O$, consists of two unique molecules linked by an $O-H\cdots N$ hydrogen bond. The conformation of both C=N bonds is *E* and the azomethine functional groups lie close to the plane of their associated benzene rings in each of the independent molecules. The dihedral angles between the two benzene rings are 83.14 (4) and 75.45 (4)°. The plane of the one of the N(CH₃)₂ units is twisted away from the benzene ring by 18.8 (2)°, indicating loss of conjugation between the lone electron pair and the benzene ring. In the crystal structure, $O-H\cdots N$ hydrogen bonds together with $C-H\cdots O$ hydrogen bonds link neighbouring supramolecular dimers into a three-dimensional network.

1. Chemical context

Schiff bases play important roles in the development of coordination chemistry related to catalysis, enzymatic reactions, and supramolecular architectures. Crystal structures of Schiff bases derived from substituted benzaldehydes and 1,3-diaminopropan-2-ol have been reported earlier (Azam, Warad, Al-Resayes *et al.*, 2012; Azam, Hussain *et al.*, 2012; Rivera *et al.*, 2016b, 2017; Elmali, 2000). The title compound, (I), acts as an important raw material for the synthesis of Schiff base complexes. As an extension of our work on the synthesis and structural characterization of such Schiff base compounds, the crystal structure of the title compound is reported here.



2. Structural commentary

The title compound crystallizes with two unique molecules in the asymmetric unit. The conformers, labeled A and B, are shown in Fig. 1. Each molecule comprises a 1,3-diamino-2hydroxypropane bridge symmetrically substituted at the 1 and 3 positions by 4-(dimethylamino)phenyl]methylidene units. The conformational differences between the two molecules are extremely small, resulting in a superstructural motif. The two molecules are related by translation along the *a*-axis





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Figure 1

The structure of the independent molecules A and B, showing the atomlabelling scheme. Displacement ellipsoids are drawn at the 50% probability level for non-H atoms.

direction. A structural overlay of the two independent molecules (r.m.s. deviation for fitting all non-H atoms = 0.097 Å) is shown in Fig. 2. The disposition of the residues attached to the N2A and N2 positions can be described by the torsion angles N2A-C5A-C51A-C56A [-9.9 (11) in molecule A] and N2-C5-C51-C56 [-14.9 (11)° in molecule B]. The two outer aromatic rings (C11-C16 and C51-C56) are inclined to one another by 83.14 (4)° in molecule A and 75.45 (4)° in molecule B.

Bond distances and angles in the benzene rings are not unusual and compare well, both between the two independent molecules and with those observed in related systems (see for example: Rivera et al., 2016b). The values for the azomethine C=N bond distances in the two molecules [1.275 (8) and 1.272 (8) in molecule A and 1.271 (8) and 1.269 (8) Å in molecule B] and the corresponding internal angles at the nitrogen atom [C1A-N1A-C2A = 117.7 (6)] and C5A-N2A - C4A = 117.7 (6) in molecule A and C1 - N1 - C2 =117.5 (6) and C5-N2-C4 = 117.6 (6) in molecule B] also agree with those reported in the literature for similar compounds (Rivera et al., 2016b) and are consistent with C=N double bonding. In both molecules, the azomethine groups adopt an E,E conformation, as can be seen from the torsion angles $C2A - N1A - C1A - C11A = 177.8 (6)^{\circ}$ and C4A - N2A - C5A - C51A = 179.9 (6)° in molecule A and



Figure 2

The structural overlay of the independent molecules A (green dashed) and B (purple) of the title compound.

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

$\overline{D - \mathbf{H} \cdots A}$	D-H	$H \cdot \cdot \cdot A$	$D \cdots A$	$D - \mathbf{H} \cdot \cdot \cdot A$
$O1-H1\cdots N2A^{i}$	0.84 (8)	2.06 (8)	2.889 (7)	170 (7)
$O1A - H1A \cdot \cdot \cdot N2$	0.84(8)	2.04 (8)	2.863 (7)	166 (7)
$C57 - H57B \cdots O1^{ii}$	0.98	2.53	3.171 (9)	123
$C57A - H57E \cdots O1A^{iii}$	0.98	2.36	3.211 (8)	145

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

 $C2-N1-C1-C11 = 178.6 (6)^{\circ}$ and C4-N2-C5-C51 = 177.0 (6) in molecule *B*.

The two dimethylamino substituents in molecule *B* are essentially coplanar with the benzene rings to which they are bound with torsion angles $C17-N3-C14-C13 = -3.1 (11)^{\circ}$ and $C57-N4-C54-C53 = -2.9 (11)^{\circ}$ and with dihedral angles between the NMe₂ plane and the benzene ring of 0.57 (2) and 4.60 (2)°, respectively, whilst in molecule *A* the corresponding torsional angles C17A-N3A-C14A-C13A and C57A-N4A-C54A-C53A are 2.2 (11) and 8.3 (10)°, respectively. The dihedral angles between the two dimethylamino groups (N3*A* and N4*A*) and the benzene rings are 5.09 (22) and 18.8 (2)° respectively, indicating that the lone electron pair of the N4*A* atom may not be completely conjugated with the benzene ring (C51*A*-C56*A*).

3. Supramolecular features

Through $O-H\cdots N$ hydrogen-bonding interactions [2.863 (7) Å] between O1A-H1A and the nitrogen N2 (Table 1), the two independent molecules interact to form C(5) chains running along the *a* axis (Fig. 3). The chains are linked into a three-dimensional framework by a pair of weaker intermolecular $C57-H57B\cdots O1^{ii}$ and $C57A-H57E\cdots O1A^{iii}$ hydrogen bonds (Table 1).





Crystal packing of the title compound, indicating the $O-H\cdots N$ hydrogen bonds (dashed lines), which result in chains along the *a*-axis direction.

4. Database survey

A search in the Cambridge Crystallographic Database (Groom *et al.*, 2016) for the fragment 1,3-bis[(benzyl-idene)amino]propan-2-ol yielded the following structures: N,N'-[(2-hydroxy-1,3-propanediyl)bis(nitrilomethylylidene-2,1-phenylene)]bis(4-methylbenzenesulfonamide) (Popov *et al.*, 2009), 2,2'-[(2-hydroxypropane-1,3-diyl)bis(nitrilomethylylidene)]diphenol (Azam, Hussain *et al.*, 2012), 1,3-bis(2-hydroxy-5-bromosalicylideneamine)propan-2-ol (Elmali, 2000), 1,3-bis[(*E*)-(2-chlorobenzylidene)amino]propan-2-ol (Azam, Warad, Al-Resayes *et al.*, 2012) and 1,3-bis[(4-methoxybenzylidene)amino]propan-2-ol (Rivera *et al.* 2016*b*). In each of these structures, the N=C double bonds adopt *E* conformations.

5. Synthesis and crystallization

The title compound was prepared as described by (Rivera *et al.* 2016*a*). The crude product was recrystallized from diethyl ether solution by slow evaporation of the solvent, giving colorless crystals suitable for X-ray diffraction (m.p. 396.8–398 K; yield 40%).

6. Refinement

Crystal data, data collection and structure refinement details are summarized in Table 2. The coordinates of the hydroxyl H atom were refined with $U_{iso}(H) = 1.5U_{eq}(O)$. The remaining H atoms were positioned geometrically and allowed to ride on their parent atoms, with d(C-H) = 0.95 Å for aromatic and azomethine atoms, d(C-H) = 0.98 Å for methyl, d(C-H) =0.99 Å for methylene, d(C-H) = 1.00 Å for tertiary CH. The $U_{iso}(H)$ values were constrained to $1.5U_{eq}(C_{methyl})$ or $1.2U_{eq}(C)$ for the remaining H atoms. The structure shows signs of a superstructure. The two molecules are related by a translation of 1/2 along the *a* axis. However, if the structure is refined in a cell with the *a* axis halved, the displacement parameters of one NMe₂ group and some of the C atoms of the phenyl ring to which this group is attached are significantly enlarged (Fig. 4). Shifting one molecule by $\frac{1}{2}$ in the *a*-axis



Figure 4 Perspective view of the molecule if the structure is refined in a cell with the *a* axis halved.

Table	2	
Experi	mental	details.

Crystal data	
Chemical formula	$C_{21}H_{28}N_4O$
M _r	352.47
Crystal system, space group	Monoclinic, $P2_1$
Temperature (K)	173
a, b, c (Å)	9.1456 (10), 10.5860 (8), 19.974 (2)
β (°)	97.110 (9)
$V(Å^3)$	1918.9 (3)
Ζ	4
Radiation type	Μο Κα
$\mu \text{ (mm}^{-1})$	0.08
Crystal size (mm)	$0.22 \times 0.03 \times 0.03$
Data collection	
Diffractometer	STOE IPDS II two-circle
Absorption correction	Multi-scan (X-AREA; Stoe & Cie, 2001)
T_{\min}, T_{\max}	0.426, 1.000
No. of measured, independent and observed $[I > 2\sigma(I)]$ reflections	16633, 6852, 3688
R _{int}	0.069
$(\sin \theta / \lambda)_{\rm max} ({\rm \AA}^{-1})$	0.609
Refinement	
$R[F^2 > 2\sigma(F^2)], wR(F^2), S$	0.063, 0.156, 0.93
No. of reflections	6852
No. of parameters	483
No. of restraints	1
H-atom treatment	H atoms treated by a mixture of independent and constrained refinement
$\Delta \rho_{\rm max}, \Delta \rho_{\rm min} \ ({ m e} \ { m \AA}^{-3})$	0.17, -0.22

Computer programs: X-AREA (Stoe & Cie, 2001), SHELXS (Sheldrick, 2008), SHELXL2014/7 (Sheldrick, 2015) and XP in SHELXTL-Plus (Sheldrick, 2008).

direction, it becomes obvious how similar the two molecules are. Nevertheless, there are small differences in their overall conformation (Fig. 5). As a result of that, we opted to refine the structure using the larger unit cell with two molecules in the asymmetric unit.

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Partial packing diagram of the title compound with one molecule shifted by $x = \frac{1}{2}$, y = 0, z = 0, showing similarities and differences between the two molecules.

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1,3-Bis{(*E*)-[4-(dimethylamino)benzylidene]amino}propan-2-ol: chain structure formation *via* an O—H···N hydrogen bond

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Computing details

Data collection: *X-AREA* (Stoe & Cie, 2001); cell refinement: *X-AREA* (Stoe & Cie, 2001); data reduction: *X-AREA* (Stoe & Cie, 2001); program(s) used to solve structure: *SHELXS* (Sheldrick, 2008); program(s) used to refine structure: *SHELXL2014/7* (Sheldrick, 2015); molecular graphics: *XP* in *SHELXTL-Plus* (Sheldrick, 2008); software used to prepare material for publication: *SHELXL2014/7* (Sheldrick, 2015).

F(000) = 760

 $\theta = 3.2 - 25.9^{\circ}$

 $\mu = 0.08 \text{ mm}^{-1}$

Needle, colourless

 $0.22\times0.03\times0.03~mm$

T = 173 K

 $D_{\rm x} = 1.220 {\rm Mg} {\rm m}^{-3}$

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

Cell parameters from 8852 reflections

1,3-Bis{(E)-[4-(dimethylamino)benzylidene]amino}propan-2-ol

Crystal data

 $C_{21}H_{28}N_4O$ $M_r = 352.47$ Monoclinic, $P2_1$ a = 9.1456 (10) Å b = 10.5860 (8) Å c = 19.974 (2) Å $\beta = 97.110 (9)^{\circ}$ $V = 1918.9 (3) Å^3$ Z = 4

Data collection

STOE IPDS II two-circle-	16633 measured reflections
diffractometer	6852 independent reflections
Radiation source: Genix 3D IµS microfocus X-	3688 reflections with $I > 2\sigma(I)$
ray source	$R_{\rm int} = 0.069$
ω scans	$\theta_{\text{max}} = 25.7^{\circ}, \ \theta_{\text{min}} = 3.2^{\circ}$
Absorption correction: multi-scan	$h = -11 \rightarrow 11$
(X-Area; Stoe & Cie, 2001)	$k = -11 \rightarrow 12$
$T_{\min} = 0.426, \ T_{\max} = 1.000$	$l = -24 \rightarrow 24$

Refinement

Refinement on F^2 Least-squares matrix: full $R[F^2 > 2\sigma(F^2)] = 0.063$ $wR(F^2) = 0.156$ S = 0.936852 reflections 483 parameters 1 restraint Hydrogen site location: mixed H atoms treated by a mixture of independent and constrained refinement $w = 1/[\sigma^2(F_o^2) + (0.0713P)^2]$ where $P = (F_o^2 + 2F_c^2)/3$ $(\Delta/\sigma)_{max} < 0.001$ $\Delta\rho_{max} = 0.17$ e Å⁻³ $\Delta\rho_{min} = -0.22$ e Å⁻³

Special details

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

	x	у	Ζ	$U_{ m iso}$ */ $U_{ m eq}$
01	-0.2650 (5)	0.6469 (4)	0.2463 (2)	0.0338 (11)
H1	-0.333 (9)	0.594 (7)	0.243 (4)	0.051*
N1	-0.0704 (6)	0.5677 (5)	0.4156 (3)	0.0337 (13)
N2	-0.0059 (6)	0.4851 (5)	0.2177 (3)	0.0325 (13)
N3	0.4143 (8)	0.5502 (6)	0.6804 (3)	0.056 (2)
N4	0.1885 (8)	0.4185 (7)	-0.0846 (3)	0.0525 (18)
C1	-0.0252 (7)	0.6501 (6)	0.4596 (3)	0.0318 (15)
H1B	-0.0717	0.7305	0.4578	0.038*
C2	-0.1945 (7)	0.6018 (7)	0.3656 (3)	0.0329 (15)
H2A	-0.2318	0.6862	0.3764	0.040*
H2B	-0.2752	0.5401	0.3672	0.040*
C3	-0.1482 (7)	0.6038 (6)	0.2949 (3)	0.0280 (14)
H3	-0.0654	0.6657	0.2953	0.034*
C4	-0.0905 (7)	0.4759 (6)	0.2754 (3)	0.0295 (14)
H4A	-0.0268	0.4401	0.3145	0.035*
H4B	-0.1745	0.4177	0.2640	0.035*
C5	-0.0408 (7)	0.4112 (6)	0.1683 (3)	0.0291 (14)
Н5	-0.1163	0.3509	0.1720	0.035*
C11	0.0959 (7)	0.6257 (6)	0.5131 (3)	0.0286 (14)
C12	0.1407 (8)	0.7210 (6)	0.5603 (3)	0.0349 (16)
H12	0.0966	0.8023	0.5552	0.042*
C13	0.2491 (8)	0.6979 (6)	0.6143 (3)	0.0346 (16)
H13	0.2791	0.7642	0.6451	0.041*
C14	0.3151 (8)	0.5775 (6)	0.6241 (3)	0.0340 (16)
C15	0.2724 (7)	0.4838 (6)	0.5751 (3)	0.0313 (15)
H15	0.3170	0.4027	0.5792	0.038*
C16	0.1666 (7)	0.5092 (6)	0.5215 (3)	0.0282 (14)
H16	0.1408	0.4448	0.4891	0.034*
C17	0.4596 (10)	0.6471 (7)	0.7298 (4)	0.050 (2)
H17A	0.5096	0.7152	0.7084	0.075*
H17B	0.5272	0.6106	0.7667	0.075*
H17C	0.3728	0.6810	0.7478	0.075*
C18	0.4822 (8)	0.4275 (7)	0.6890 (4)	0.0458 (18)
H18A	0.4071	0.3642	0.6956	0.069*
H18B	0.5574	0.4287	0.7285	0.069*
H18C	0.5283	0.4061	0.6487	0.069*
C51	0.0284 (8)	0.4128 (7)	0.1056 (3)	0.0330 (16)
C52	0.0002 (8)	0.3146 (7)	0.0598 (3)	0.0431 (17)
H52	-0.0583	0.2455	0.0713	0.052*

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

C53	0.0546 (9)	0.3135 (8)	-0.0025 (3)	0.0458 (19)
Н53	0.0343	0.2439	-0.0322	0.055*
C54	0.1384 (8)	0.4140 (8)	-0.0211 (4)	0.0390 (17)
C55	0.1671 (8)	0.5154 (7)	0.0254 (3)	0.0376 (17)
Н55	0.2227	0.5861	0.0138	0.045*
C56	0.1155 (8)	0.5119(7)	0.0867 (3)	0.0351 (16)
Н56	0.1396	0.5792	0.1175	0.042*
C57	0.1555 (11)	0.3106 (9)	-0.1292(4)	0.066 (3)
H57A	0.1979	0.2340	-0.1072	0.099*
H57B	0.1980	0.3246	-0.1713	0.099*
H57C	0.0485	0.3008	-0.1392	0.099*
C58	0.2838(10)	0.5179 (9)	-0.1020(4)	0.059(2)
H58A	0.2329	0.5992	-0.1009	0.089*
H58B	0.3103	0.5030	-0.1474	0.089*
H58C	0.3734	0.5193	-0.0695	0.089*
014	0.3754 0.2364 (5)	0.5175	0.0075	0.089
HIA	0.2304(3) 0.175(9)	0.0331(+) 0.598(7)	0.2491(2) 0.236(4)	0.046*
NIA	0.175(9) 0.4307(6)	0.598(7)	0.230(4) 0.4178(2)	0.040
INTA N2A	0.4307(0)	0.3729(3) 0.4805(5)	0.4178(2) 0.2205(2)	0.0321(13) 0.0272(12)
INZA	0.4641(0)	0.4803(3)	0.2203(3)	0.0272(12)
NJA NJA	0.9201(8)	0.3491(0) 0.2641(7)	0.0802(3)	0.0328(18)
N4A	0.0733(7)	0.3041(7)	-0.0780(3)	0.0495(10)
	0.4784 (8)	0.0343 (0)	0.4623 (3)	0.0320 (15)
HIAI	0.4335	0.7353	0.4603	0.038*
C2A	0.3069 (7)	0.6088 (6)	0.3686 (3)	0.0301 (14)
H2A1	0.2727	0.6943	0.3792	0.036*
H2A2	0.2244	0.5490	0.3710	0.036*
C3A	0.3525 (7)	0.6076 (6)	0.2969 (3)	0.0267 (14)
H3A	0.4386	0.6658	0.2967	0.032*
C4A	0.4027 (8)	0.4766 (6)	0.2788 (3)	0.0314 (15)
H4A1	0.4663	0.4404	0.3178	0.038*
H4A2	0.3156	0.4210	0.2688	0.038*
C5A	0.4427 (8)	0.4067 (6)	0.1714 (3)	0.0295 (15)
H5A	0.3609	0.3534	0.1759	0.035*
C11A	0.5983 (7)	0.6298 (6)	0.5160 (3)	0.0261 (14)
C12A	0.6474 (7)	0.7222 (6)	0.5632 (3)	0.0317 (15)
H12A	0.6067	0.8047	0.5582	0.038*
C13A	0.7528 (8)	0.6973 (6)	0.6165 (3)	0.0345 (16)
H13A	0.7825	0.7622	0.6481	0.041*
C14A	0.8175 (8)	0.5775 (6)	0.6252 (3)	0.0336 (15)
C15A	0.7757 (8)	0.4861 (6)	0.5758 (3)	0.0355 (16)
H15A	0.8212	0.4053	0.5791	0.043*
C16A	0.6686 (7)	0.5124 (6)	0.5223 (3)	0.0295 (14)
H16A	0.6423	0.4492	0.4892	0.035*
C17A	0.9607 (10)	0.6408 (7)	0.7321 (4)	0.052 (2)
H17D	1.0069	0.7135	0.7128	0.077*
H17E	1.0305	0.6029	0.7676	0.077*
H17F	0.8725	0.6686	0.7512	0.077*
C18A	0.9840 (8)	0.4245 (7)	0.6888 (4)	0.0457 (18)

H18D	0.9081	0.3642	0.6985	0.069*	
H18E	1.0637	0.4255	0.7264	0.069*	
H18F	1.0235	0.3994	0.6474	0.069*	
C51A	0.5096 (8)	0.3966 (6)	0.1087 (3)	0.0304 (16)	
C52A	0.4621 (8)	0.3029 (6)	0.0628 (3)	0.0386 (16)	
H52A	0.3901	0.2446	0.0740	0.046*	
C53A	0.5148 (9)	0.2908 (7)	0.0017 (4)	0.0457 (18)	
H53A	0.4794	0.2247	-0.0282	0.055*	
C54A	0.6205 (9)	0.3753 (8)	-0.0169 (3)	0.0409 (19)	
C55A	0.6708 (8)	0.4707 (8)	0.0289 (3)	0.0403 (17)	
H55A	0.7434	0.5288	0.0180	0.048*	
C56A	0.6136 (8)	0.4806 (7)	0.0914 (3)	0.0371 (17)	
H56A	0.6475	0.5462	0.1219	0.045*	
C57A	0.6067 (10)	0.2760 (8)	-0.1289 (3)	0.063 (3)	
H57D	0.6115	0.1904	-0.1100	0.095*	
H57E	0.6600	0.2786	-0.1685	0.095*	
H57F	0.5034	0.2993	-0.1421	0.095*	
C58A	0.7565 (10)	0.4673 (10)	-0.1032 (4)	0.065 (3)	
H58D	0.6943	0.5429	-0.1090	0.097*	
H58E	0.7881	0.4437	-0.1466	0.097*	
H58F	0.8433	0.4850	-0.0706	0.097*	

Atomic displacement parameters $(Å^2)$

	U^{11}	U ²²	U^{33}	U^{12}	U^{13}	U^{23}
01	0.038 (3)	0.032 (3)	0.031 (2)	-0.002 (2)	-0.002 (2)	0.0081 (19)
N1	0.038 (3)	0.036 (3)	0.028 (3)	0.002 (2)	0.009 (3)	-0.001 (2)
N2	0.034 (3)	0.034 (3)	0.030 (3)	0.006 (2)	0.006 (2)	0.001 (2)
N3	0.081 (5)	0.032 (4)	0.047 (4)	-0.005 (3)	-0.027 (4)	0.001 (3)
N4	0.051 (4)	0.078 (5)	0.030 (3)	0.005 (4)	0.009 (3)	-0.013 (3)
C1	0.029 (4)	0.032 (4)	0.035 (4)	0.009 (3)	0.007 (3)	0.000 (3)
C2	0.028 (4)	0.041 (4)	0.031 (3)	0.003 (3)	0.007 (3)	0.001 (3)
C3	0.028 (3)	0.026 (3)	0.030 (3)	0.001 (3)	0.005 (3)	0.000(2)
C4	0.030 (3)	0.032 (3)	0.027 (3)	-0.005 (3)	0.004 (3)	0.006 (3)
C5	0.027 (3)	0.028 (3)	0.031 (3)	-0.002 (3)	-0.002 (3)	-0.002 (3)
C11	0.026 (3)	0.032 (3)	0.030(3)	-0.002 (3)	0.012 (3)	-0.001 (3)
C12	0.043 (4)	0.025 (3)	0.036 (4)	0.001 (3)	0.002 (3)	0.002 (3)
C13	0.044 (4)	0.026 (3)	0.034 (4)	-0.008 (3)	0.006 (3)	-0.008 (3)
C14	0.040 (4)	0.033 (4)	0.028 (3)	-0.009 (3)	0.001 (3)	0.008 (3)
C15	0.036 (4)	0.024 (3)	0.033 (3)	-0.003 (3)	0.003 (3)	0.001 (2)
C16	0.027 (3)	0.027 (3)	0.031 (3)	-0.005 (3)	0.006 (3)	-0.004 (2)
C17	0.064 (5)	0.046 (4)	0.037 (4)	-0.012 (4)	-0.010 (4)	-0.002 (3)
C18	0.054 (5)	0.043 (4)	0.037 (4)	-0.004(4)	-0.005 (3)	0.009 (3)
C51	0.038 (4)	0.035 (4)	0.025 (3)	0.013 (3)	0.002 (3)	-0.005 (3)
C52	0.050 (4)	0.044 (4)	0.035 (4)	0.004 (3)	0.003 (3)	0.001 (3)
C53	0.054 (4)	0.052 (4)	0.030 (3)	0.009 (4)	0.000 (3)	-0.018 (3)
C54	0.033 (4)	0.052 (4)	0.032 (3)	0.007 (3)	0.001 (3)	-0.004 (3)
C55	0.038 (4)	0.047 (4)	0.027 (3)	0.001 (3)	-0.001 (3)	-0.004 (3)

C56	0.031 (4)	0.044 (4)	0.031 (3)	-0.004 (3)	0.003 (3)	-0.008 (3)
C57	0.086 (6)	0.073 (6)	0.038 (4)	0.029 (5)	0.008 (4)	-0.017 (4)
C58	0.054 (5)	0.088 (6)	0.041 (4)	0.015 (4)	0.030 (4)	0.012 (4)
O1A	0.031 (3)	0.035 (3)	0.026 (2)	0.000 (2)	0.0010 (19)	0.0069 (18)
N1A	0.032 (3)	0.038 (3)	0.026 (3)	0.002 (2)	0.003 (2)	-0.003 (2)
N2A	0.028 (3)	0.025 (3)	0.027 (3)	0.003 (2)	0.001 (2)	-0.002 (2)
N3A	0.068 (5)	0.037 (4)	0.044 (3)	-0.004 (3)	-0.030 (3)	0.003 (3)
N4A	0.046 (4)	0.074 (4)	0.029 (3)	0.004 (3)	0.011 (3)	-0.012 (3)
C1A	0.037 (4)	0.033 (4)	0.027 (4)	0.002 (3)	0.007 (3)	-0.005 (3)
C2A	0.032 (4)	0.033 (4)	0.025 (3)	-0.002 (3)	0.007 (3)	-0.001 (3)
C3A	0.029 (3)	0.027 (3)	0.025 (3)	-0.003 (3)	0.005 (3)	0.000(2)
C4A	0.039 (4)	0.029 (3)	0.026 (3)	0.005 (3)	0.005 (3)	0.003 (3)
C5A	0.032 (4)	0.023 (3)	0.032 (3)	0.006 (3)	0.001 (3)	0.006 (2)
C11A	0.028 (3)	0.027 (3)	0.023 (3)	-0.001 (3)	0.004 (3)	0.000(2)
C12A	0.037 (4)	0.025 (3)	0.034 (3)	0.003 (3)	0.004 (3)	-0.005 (3)
C13A	0.040 (4)	0.035 (4)	0.027 (3)	-0.005 (3)	-0.002 (3)	-0.002 (3)
C14A	0.031 (4)	0.032 (4)	0.036 (4)	-0.002 (3)	-0.003 (3)	0.000 (3)
C15A	0.045 (4)	0.024 (3)	0.037 (4)	-0.004 (3)	0.004 (3)	0.003 (3)
C16A	0.035 (4)	0.027 (3)	0.026 (3)	-0.008 (3)	0.004 (3)	-0.001 (2)
C17A	0.064 (5)	0.044 (4)	0.040 (4)	-0.007 (4)	-0.021 (4)	-0.003 (3)
C18A	0.046 (4)	0.040 (4)	0.047 (4)	-0.002 (4)	-0.014 (3)	0.015 (3)
C51A	0.027 (3)	0.033 (4)	0.029 (3)	0.006 (3)	-0.004 (3)	-0.003 (2)
C52A	0.052 (4)	0.031 (3)	0.031 (3)	0.005 (3)	0.003 (3)	-0.006 (3)
C53A	0.060 (5)	0.042 (4)	0.034 (3)	0.008 (4)	0.001 (3)	-0.008 (3)
C54A	0.036 (4)	0.057 (5)	0.028 (3)	0.024 (4)	-0.005 (3)	-0.014 (3)
C55A	0.027 (3)	0.065 (5)	0.029 (3)	0.001 (3)	0.006 (3)	-0.007 (3)
C56A	0.032 (4)	0.053 (4)	0.026 (3)	0.002 (3)	0.001 (3)	-0.008 (3)
C57A	0.090 (6)	0.074 (6)	0.024 (3)	0.038 (5)	0.001 (4)	-0.017 (3)
C58A	0.059 (5)	0.097 (7)	0.037 (4)	0.021 (5)	0.005 (4)	0.001 (4)

Geometric parameters (Å, °)

01—C3	1.426 (7)	O1A—C3A	1.429 (7)
O1—H1	0.84 (8)	O1A—H1A	0.84 (8)
N1-C1	1.271 (8)	N1A—C1A	1.275 (8)
N1—C2	1.461 (8)	N1A—C2A	1.454 (8)
N2—C5	1.269 (8)	N2A—C5A	1.272 (8)
N2-C4	1.470 (8)	N2A—C4A	1.460 (8)
N3—C14	1.386 (9)	N3A—C14A	1.388 (8)
N3—C18	1.440 (10)	N3A—C17A	1.434 (9)
N3—C17	1.448 (9)	N3A—C18A	1.444 (10)
N4—C54	1.400 (9)	N4A—C54A	1.383 (9)
N4—C58	1.436 (11)	N4A—C57A	1.449 (10)
N4—C57	1.458 (11)	N4A—C58A	1.452 (12)
C1-C11	1.463 (9)	C1A—C11A	1.459 (8)
C1—H1B	0.9500	C1A—H1A1	0.9500
C2—C3	1.523 (9)	C2A—C3A	1.542 (8)
C2—H2A	0.9900	C2A—H2A1	0.9900

C2—H2B	0.9900	C2A—H2A2	0.9900
C3—C4	1.521 (9)	C3A—C4A	1.518 (8)
С3—Н3	1.0000	СЗА—НЗА	1.0000
C4—H4A	0.9900	C4A—H4A1	0.9900
C4—H4B	0.9900	C4A—H4A2	0.9900
C5—C51	1.472 (10)	C5A—C51A	1.465 (10)
С5—Н5	0.9500	C5A—H5A	0.9500
C11—C16	1.393 (9)	C11A—C12A	1.394 (9)
C11—C12	1.407 (9)	C11A—C16A	1.397 (9)
C12—C13	1.395 (10)	C12A—C13A	1.371 (9)
С12—Н12	0.9500	C12A—H12A	0.9500
C13—C14	1.413 (10)	C13A—C14A	1.400 (9)
С13—Н13	0.9500	C13A—H13A	0.9500
C14—C15	1.415 (9)	C14A—C15A	1.400 (9)
C15—C16	1.377 (9)	C15A—C16A	1.386 (9)
С15—Н15	0.9500	C15A—H15A	0.9500
C16—H16	0.9500	C16A—H16A	0.9500
C17—H17A	0.9800	C17A—H17D	0.9800
C17—H17B	0.9800	C17A - H17E	0.9800
C17_H17C	0.9800	C17A H17E	0.9800
C18—H18A	0.9800	C184—H18D	0.9800
C18 H18B	0.9800		0.9800
	0.9800	C18A H18E	0.9800
$C_{10} - 1118C$	1.387(10)	C_{10A} C_{50A}	1.377(10)
C51_C52	1.307(10) 1.307(10)	C_{51A} C_{52A}	1.377(10) 1.384(0)
$C_{51} = C_{50}$	1.397(10) 1.207(10)	C51A = C52A	1.364(9) 1.272(10)
C52—C55	1.397 (10)	C52A—C55A	1.373 (10)
C52—H52	0.9500	C52A—H52A	0.9500
C53-C54	1.388 (11)	C53A = C54A	1.399 (12)
С53—Н53	0.9500	C53A—H53A	0.9500
C34—C33	1.423 (10)	C54A—C55A	1.401 (10)
055-056	1.365 (10)	C55A—C56A	1.416 (10)
C55—H55	0.9500	C55A—H55A	0.9500
C56—H56	0.9500	С56А—Н56А	0.9500
С57—Н57А	0.9800	C57A—H57D	0.9800
С57—Н57В	0.9800	C57A—H57E	0.9800
С57—Н57С	0.9800	C57A—H57F	0.9800
С58—Н58А	0.9800	C58A—H58D	0.9800
C58—H58B	0.9800	C58A—H58E	0.9800
C58—H58C	0.9800	C58A—H58F	0.9800
C3—O1—H1	109 (5)	C3A—O1A—H1A	112 (5)
C1—N1—C2	117.5 (6)	C1A—N1A—C2A	117.7 (6)
C5—N2—C4	117.6 (6)	C5A—N2A—C4A	117.7 (6)
C14—N3—C18	121.0 (6)	C14A—N3A—C17A	121.0 (6)
C14—N3—C17	120.6 (6)	C14A—N3A—C18A	121.1 (6)
C18—N3—C17	118.3 (6)	C17A - N3A - C18A	117.9 (6)
C54—N4—C58	121.4 (7)	C54A—N4A—C57A	120.6 (7)
C54—N4—C57	117.5 (7)	C54A—N4A—C58A	119.6 (7)
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C58—N4—C57	120.7 (7)	C57A—N4A—C58A	116.2 (6)
N1—C1—C11	122.4 (6)	N1A—C1A—C11A	123.6 (6)
N1—C1—H1B	118.8	N1A—C1A—H1A1	118.2
C11—C1—H1B	118.8	C11A—C1A—H1A1	118.2
N1—C2—C3	110.8 (5)	N1A—C2A—C3A	110.4 (5)
N1—C2—H2A	109.5	N1A—C2A—H2A1	109.6
C3—C2—H2A	109.5	C3A—C2A—H2A1	109.6
N1—C2—H2B	109.5	N1A—C2A—H2A2	109.6
С3—С2—Н2В	109.5	C3A—C2A—H2A2	109.6
H2A—C2—H2B	108.1	H2A1—C2A—H2A2	108.1
O1—C3—C4	111.4 (5)	O1A—C3A—C4A	112.6 (5)
O1—C3—C2	111.3 (5)	O1A—C3A—C2A	110.8 (5)
C4—C3—C2	111.7 (5)	C4A—C3A—C2A	110.6 (5)
O1—C3—H3	107.4	O1A—C3A—H3A	107.5
С4—С3—Н3	107.4	С4А—С3А—НЗА	107.5
С2—С3—Н3	107.4	С2А—С3А—НЗА	107.5
N2—C4—C3	112.0 (5)	N2A—C4A—C3A	111.3 (5)
N2—C4—H4A	109.2	N2A—C4A—H4A1	109.4
C3—C4—H4A	109.2	C3A—C4A—H4A1	109.4
N2—C4—H4B	109.2	N2A—C4A—H4A2	109.4
C3—C4—H4B	109.2	C3A—C4A—H4A2	109.4
H4A—C4—H4B	107.9	H4A1—C4A—H4A2	108.0
N2—C5—C51	124.1 (6)	N2A—C5A—C51A	126.0 (7)
N2—C5—H5	117.9	N2A—C5A—H5A	117.0
С51—С5—Н5	117.9	С51А—С5А—Н5А	117.0
C16—C11—C12	117.5 (6)	C12A—C11A—C16A	117.2 (6)
C16—C11—C1	122.9 (6)	C12A—C11A—C1A	121.5 (6)
C12—C11—C1	119.6 (6)	C16A—C11A—C1A	121.3 (6)
C13—C12—C11	120.9 (6)	C13A—C12A—C11A	121.8 (6)
C13—C12—H12	119.6	C13A—C12A—H12A	119.1
C11—C12—H12	119.6	C11A—C12A—H12A	119.1
C12—C13—C14	121.0 (6)	C12A—C13A—C14A	121.0 (6)
C12—C13—H13	119.5	C12A—C13A—H13A	119.5
C14—C13—H13	119.5	C14A—C13A—H13A	119.5
N3—C14—C13	121.7 (6)	N3A—C14A—C15A	120.5 (6)
N3—C14—C15	120.8 (6)	N3A—C14A—C13A	121.9 (6)
C13—C14—C15	117.4 (6)	C15A—C14A—C13A	117.7 (6)
C16—C15—C14	120.5 (6)	C16A—C15A—C14A	120.6 (6)
C16—C15—H15	119.7	C16A—C15A—H15A	119.7
C14—C15—H15	119.7	C14A—C15A—H15A	119.7
C15—C16—C11	122.5 (6)	C15A—C16A—C11A	121.4 (6)
C15—C16—H16	118.7	C15A—C16A—H16A	119.3
C11—C16—H16	118.7	C11A—C16A—H16A	119.3
N3—C17—H17A	109.5	N3A—C17A—H17D	109.5
N3—C17—H17B	109.5	N3A—C17A—H17E	109.5
H17A—C17—H17B	109.5	H17D—C17A—H17E	109.5
N3—C17—H17C	109.5	N3A—C17A—H17F	109.5
H17A—C17—H17C	109.5	H17D—C17A—H17F	109.5

H17B—C17—H17C	109.5	H17E—C17A—H17F	109.5
N3—C18—H18A	109.5	N3A—C18A—H18D	109.5
N3—C18—H18B	109.5	N3A—C18A—H18E	109.5
H18A—C18—H18B	109.5	H18D—C18A—H18E	109.5
N3—C18—H18C	109.5	N3A—C18A—H18F	109.5
H18A—C18—H18C	109.5	H18D—C18A—H18F	109.5
H18B—C18—H18C	109.5	H18E—C18A—H18F	109.5
C52—C51—C56	116.8 (6)	C56A—C51A—C52A	117.8 (7)
C52—C51—C5	119.1 (7)	C56A—C51A—C5A	122.8 (6)
C56—C51—C5	124.0 (6)	C52A—C51A—C5A	119.3 (7)
C51—C52—C53	122.3 (8)	C53A—C52A—C51A	122.7 (7)
С51—С52—Н52	118.8	С53А—С52А—Н52А	118.6
С53—С52—Н52	118.8	С51А—С52А—Н52А	118.7
C54—C53—C52	120.1 (7)	C52A—C53A—C54A	120.4 (7)
С54—С53—Н53	119.9	С52А—С53А—Н53А	119.8
С52—С53—Н53	119.9	С54А—С53А—Н53А	119.8
C53—C54—N4	121.4 (7)	N4A—C54A—C53A	120.7 (7)
C53—C54—C55	117.8 (7)	N4A—C54A—C55A	121.3 (8)
N4—C54—C55	120.7 (7)	C53A—C54A—C55A	118.0 (7)
C56—C55—C54	120.5 (7)	C54A—C55A—C56A	120.1 (7)
С56—С55—Н55	119.7	C54A—C55A—H55A	120.0
С54—С55—Н55	119.7	C56A—C55A—H55A	120.0
C55—C56—C51	122.3 (7)	C51A—C56A—C55A	121.0 (7)
С55—С56—Н56	118.8	C51A—C56A—H56A	119.5
С51—С56—Н56	118.8	C55A—C56A—H56A	119.5
N4—C57—H57A	109.5	N4A—C57A—H57D	109.5
N4—C57—H57B	109.5	N4A—C57A—H57E	109.5
H57A—C57—H57B	109.5	H57D—C57A—H57E	109.5
N4—C57—H57C	109.5	N4A—C57A—H57F	109.5
Н57А—С57—Н57С	109.5	H57D—C57A—H57F	109.5
Н57В—С57—Н57С	109.5	H57E—C57A—H57F	109.5
N4—C58—H58A	109.5	N4A—C58A—H58D	109.5
N4—C58—H58B	109.5	N4A—C58A—H58E	109.5
H58A—C58—H58B	109.5	H58D—C58A—H58E	109.5
N4—C58—H58C	109.5	N4A—C58A—H58F	109.5
H58A—C58—H58C	109.5	H58D—C58A—H58F	109.5
H58B—C58—H58C	109.5	H58E—C58A—H58F	109.5
C2—N1—C1—C11	178.6 (6)	C2A—N1A—C1A—C11A	177.8 (6)
C1—N1—C2—C3	116.0 (7)	C1A—N1A—C2A—C3A	117.4 (6)
N1—C2—C3—O1	-175.3(5)	N1A—C2A—C3A—O1A	-174.1(5)
N1-C2-C3-C4	59.4 (8)	N1A—C2A—C3A—C4A	60.4 (7)
C5-N2-C4-C3	-128.6(6)	C5A - N2A - C4A - C3A	-127.7(6)
01 - C3 - C4 - N2	71.8 (7)	O1A— $C3A$ — $C4A$ — $N2A$	71.5 (7)
$C_2 - C_3 - C_4 - N_2$	-163.0(6)	C2A - C3A - C4A - N2A	-163.9(5)
C4-N2-C5-C51	177.0 (6)	C4A - N2A - C5A - C51A	179.9 (6)
N1-C1-C11-C16	-2.8(10)	N1A-C1A-C11A-C12A	179.6 (7)
N1-C1-C11-C12	1800(7)	N1A-C1A-C11A-C16A	-0.6(10)
	100.0(7)		0.0 (10)

C16—C11—C12—C13	-1.7 (10)	C16A—C11A—C12A—C13A	-4.5 (10)
C1-C11-C12-C13	175.7 (6)	C1A—C11A—C12A—C13A	175.3 (6)
C11—C12—C13—C14	-1.2 (11)	C11A—C12A—C13A—C14A	1.0 (11)
C18—N3—C14—C13	-179.0 (7)	C17A—N3A—C14A—C15A	-178.2 (8)
C17—N3—C14—C13	-3.1 (11)	C18A—N3A—C14A—C15A	-1.3 (11)
C18—N3—C14—C15	3.1 (11)	C17A—N3A—C14A—C13A	2.2 (11)
C17—N3—C14—C15	178.9 (7)	C18A—N3A—C14A—C13A	179.1 (7)
C12-C13-C14-N3	-174.8 (7)	C12A—C13A—C14A—N3A	-177.5 (7)
C12—C13—C14—C15	3.2 (10)	C12A—C13A—C14A—C15A	2.9 (10)
N3-C14-C15-C16	175.8 (7)	N3A—C14A—C15A—C16A	177.2 (7)
C13—C14—C15—C16	-2.2 (10)	C13A—C14A—C15A—C16A	-3.1 (10)
C14-C15-C16-C11	-0.7 (10)	C14A—C15A—C16A—C11A	-0.4 (10)
C12—C11—C16—C15	2.7 (10)	C12A—C11A—C16A—C15A	4.2 (10)
C1-C11-C16-C15	-174.6 (6)	C1A—C11A—C16A—C15A	-175.6 (6)
N2-C5-C51-C52	169.0 (7)	N2A—C5A—C51A—C56A	-9.9 (11)
N2-C5-C51-C56	-14.9 (11)	N2A—C5A—C51A—C52A	173.4 (7)
C56—C51—C52—C53	-0.3 (11)	C56A—C51A—C52A—C53A	0.3 (10)
C5—C51—C52—C53	176.0 (7)	C5A—C51A—C52A—C53A	177.1 (7)
C51—C52—C53—C54	-1.0 (12)	C51A—C52A—C53A—C54A	-0.3 (11)
C52—C53—C54—N4	-176.9 (7)	C57A—N4A—C54A—C53A	8.3 (10)
C52—C53—C54—C55	0.7 (11)	C58A—N4A—C54A—C53A	166.3 (7)
C58—N4—C54—C53	-175.5 (7)	C57A—N4A—C54A—C55A	-171.8 (7)
C57—N4—C54—C53	-2.9 (11)	C58A—N4A—C54A—C55A	-13.9 (11)
C58—N4—C54—C55	7.0 (12)	C52A—C53A—C54A—N4A	-179.6 (7)
C57—N4—C54—C55	179.6 (7)	C52A—C53A—C54A—C55A	0.6 (11)
C53—C54—C55—C56	1.1 (11)	N4A—C54A—C55A—C56A	179.4 (6)
N4—C54—C55—C56	178.6 (7)	C53A—C54A—C55A—C56A	-0.8 (11)
C54—C55—C56—C51	-2.5 (12)	C52A—C51A—C56A—C55A	-0.5 (10)
C52—C51—C56—C55	2.1 (11)	C5A—C51A—C56A—C55A	-177.2 (6)
C5-C51-C56-C55	-174.0 (7)	C54A—C55A—C56A—C51A	0.8 (11)

Hydrogen-bond geometry (Å, °)

<i>D</i> —Н	H···A	$D \cdots A$	<i>D</i> —H… <i>A</i>
0.84 (8)	2.06 (8)	2.889 (7)	170 (7)
0.84 (8)	2.04 (8)	2.863 (7)	166 (7)
0.98	2.53	3.171 (9)	123
0.98	2.36	3.211 (8)	145
	<i>D</i> —H 0.84 (8) 0.84 (8) 0.98 0.98	D—H H···A 0.84 (8) 2.06 (8) 0.84 (8) 2.04 (8) 0.98 2.53 0.98 2.36	DHH···AD···A0.84 (8)2.06 (8)2.889 (7)0.84 (8)2.04 (8)2.863 (7)0.982.533.171 (9)0.982.363.211 (8)

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