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(E)-N-[(E)-3-(4-Nitrophenyl)allylidene]naphthalen-1-amine

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Key indicators: single-crystal X-ray study; T = 296 K; mean σ (C–C) = 0.002 Å; R factor = 0.044; wR factor = 0.125; data-to-parameter ratio = 15.6.

In the title compound, $C_{19}H_{14}N_2O_2$, the dihedral angle between the mean planes of the 4-nitrophenyl ring and the naphthalene ring system is 12.79 (2)°. The imine group displays a C-C-N=C torsion angle of 41.0 (2)° and the C=N group has an *E* conformation. In the crystal, weak C-H···O hydrogen bonds link molecules into layers parallel to the *b* axis.

Related literature

For the synthesis and biological activity of naphthalene compounds, see: Upadhayaya *et al.* (2010); Rokade & Sayyed (2009). For a related structure, see: Yang *et al.* (2012).



Experimental

Crystal data

Data collection

Rigaku R-AXIS RAPID diffractometer Absorption correction: multi-scan (*ABSCOR*; Rigaku, 1995) $T_{\rm min} = 0.714, T_{\rm max} = 0.991$

Refinement

$R[F^2 > 2\sigma(F^2)] = 0.044$	
$vR(F^2) = 0.125$	
S = 1.00	
3426 reflections	
220 parameters	

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

, , ,				
$D - H \cdots A$	D-H	$H \cdot \cdot \cdot A$	$D \cdots A$	$D - \mathbf{H} \cdot \cdot \cdot A$
$C5-H5\cdotsO1^{i}$ $C15-H15\cdotsO2^{ii}$	0.93 0.93	2.66 2.46	3.422 (3) 3.326 (3)	139 155

14376 measured reflections

 $R_{\rm int} = 0.038$

refinement $\Delta \rho_{\text{max}} = 0.16 \text{ e } \text{ Å}^{-3}$ $\Delta \rho_{\text{min}} = -0.24 \text{ e } \text{ Å}^{-3}$

3426 independent reflections 1824 reflections with $F^2 > 2\sigma(F^2)$

H atoms treated by a mixture of

independent and constrained

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

Data collection: *RAPID-AUTO* (Rigaku, 2006); cell refinement: *RAPID-AUTO*; data reduction: *RAPID-AUTO*; program(s) used to solve structure: *Il Milione* (Burla *et al.*, 2007); program(s) used to refine structure: *SHELXL97* (Sheldrick, 2008); molecular graphics: *CrystalStructure* (Rigaku, 2010); software used to prepare material for publication: *CrystalStructure*.

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Supplementary data and figures for this paper are available from the IUCr electronic archives (Reference: FF2100).

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(E)-N-[(E)-3-(4-Nitrophenyl)allylidene]naphthalen-1-amine

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S1. Comment

Naphthalene derivatives has been identified as new range of potent antimicrobials effective against wide range of human pathogens. They occupy a central place among medicinally important compounds due to their diverse and interesting antibiotic properties with minimum toxicity. (Rokade & Sayyed, 2009; Upadhayaya *et al.* 2010). In this paper, the title compound was synthesized and characterized by X-ray diffraction. Crystal structure of a similar compound, *N*-(Naphthalen-1-yl­methyl­idene)-4*H*-1,2,4-triazol-4-amine, was described previously by Yang *et al.* (2012).

In the title compound (Fig. 1), $C_{19}H_{14}N_2O_2$, the dihedral angle between the mean planes of the 4-nitrophenyl ring (C14— C19) and the naphthalene ring (C1—C10) is 12.79 (2)°. The imine group displays a torsion angle [C1–C10–N1–C11 = 41.0 (2)°] and the imine [C11 = N1] group has an (*E*) configuration. In the crystal, weak intermolecular C—H…O hydrogen bonds link molecules into layers parallel to the *b* axis.

S2. Experimental

To a solution of 1-naphthylamine (2.0 mmol) in anhydrous ethanol (40 ml) was treated with equimolar quantities of substituted 4-nitrocinnamaldehydes. The mixture was refluxed for 3 days, and the progress of reaction was monitored by TLC. After completion of reaction, the solvent was removed under reduced pressure. The residue was purified by flash column chromatography to afford the title compound as a yellow solid in yield 88%. Single crystals suitable for X-ray diffraction were prepared by slow evaporation of a solution of the title compound in ethanol at room temperature.

S3. Refinement

All hydrogen atoms were positioned geometrically and refined using a riding model with C—H = 0.93-0.98 Å and Uiso(H) = 1.2 or 1.5 Ueq(C).



Figure 1

The molecular structure of the title compound showing the atomic numbering and 50% probability displacement ellipsoid.

(E)-N-[(E)-3-(4-Nitrophenyl)allylidene]naphthalen-1-amine

Crystal data	
$C_{19}H_{14}N_2O_2$	V = 1507.8 (2) Å ³
$M_r = 302.33$	Z = 4
Monoclinic, $P2_1/c$	F(000) = 632.00
Hall symbol: -P 2ybc	$D_{\rm x} = 1.332 {\rm ~Mg} {\rm ~m}^{-3}$
a = 7.7021 (5) Å	Mo <i>K</i> α radiation, $\lambda = 0.71075$ Å
b = 13.8713 (12) Å	Cell parameters from 7945 reflections
c = 14.2554 (10) Å	$\theta = 3.1 - 27.5^{\circ}$
$\beta = 98.096 \ (2)^{\circ}$	$\mu=0.09~\mathrm{mm^{-1}}$

T = 296 K Chunk, yellow	$0.20 \times 0.20 \times 0.10 \text{ mm}$
Data collection	
Rigaku R-AXIS RAPID diffractometer Detector resolution: 10.000 pixels mm ⁻¹ ω scans Absorption correction: multi-scan (<i>ABSCOR</i> ; Rigaku, 1995) $T_{min} = 0.714, T_{max} = 0.991$ 14376 measured reflections	3426 independent reflections 1824 reflections with $F^2 > 2\sigma(F^2)$ $R_{int} = 0.038$ $\theta_{max} = 27.5^{\circ}$ $h = -8 \rightarrow 9$ $k = -18 \rightarrow 17$ $l = -18 \rightarrow 18$
Refinement	
Refinement on F^2 $R[F^2 > 2\sigma(F^2)] = 0.044$ $wR(F^2) = 0.125$ S = 1.00 3426 reflections 220 parameters 0 restraints Primary atom site location: structure-invariant direct methods	Secondary atom site location: difference Fourier map Hydrogen site location: inferred from neighbouring sites H atoms treated by a mixture of independent and constrained refinement $w = 1/[\sigma^2(F_o^2) + (0.0632P)^2]$ where $P = (F_o^2 + 2F_c^2)/3$ $(\Delta/\sigma)_{max} < 0.001$ $\Delta\rho_{max} = 0.16$ e Å ⁻³ $\Lambda\rho_{min} = -0.24$ e Å ⁻³

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. **Refinement**. Refinement was performed using all reflections. The weighted *R*-factor (*wR*) and goodness of fit (*S*) are based on F^2 . *R*-factor (gt) are based on *F*. The threshold expression of $F^2 > 2.0 \sigma(F^2)$ is used only for calculating *R*-factor (gt).

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

	x	У	Ζ	$U_{ m iso}$ */ $U_{ m eq}$	
01	0.83231 (18)	0.69908 (11)	0.01889 (9)	0.0893 (5)	
O2	1.04049 (17)	0.66549 (11)	0.13076 (10)	0.0912 (5)	
N1	0.28674 (17)	0.58150 (9)	0.58185 (9)	0.0559 (4)	
N2	0.8861 (2)	0.68108 (10)	0.10197 (11)	0.0639 (4)	
C1	-0.0251 (2)	0.57846 (11)	0.60108 (11)	0.0571 (4)	
C2	-0.1560 (2)	0.57496 (12)	0.65987 (13)	0.0646 (5)	
C3	-0.1124 (3)	0.57509 (12)	0.75554 (13)	0.0650 (5)	
C4	0.0649 (2)	0.57721 (10)	0.79753 (11)	0.0531 (4)	
C5	0.1143 (3)	0.57601 (12)	0.89707 (12)	0.0656 (5)	
C6	0.2852 (3)	0.57557 (13)	0.93612 (12)	0.0708 (5)	
C7	0.4176 (3)	0.57612 (12)	0.87819 (12)	0.0667 (5)	
C8	0.3764 (2)	0.57776 (11)	0.78197 (11)	0.0566 (4)	
C9	0.19908 (19)	0.57918 (10)	0.73900 (10)	0.0478 (4)	
C10	0.1495 (2)	0.58192 (10)	0.63815 (10)	0.0493 (4)	
C11	0.2713 (3)	0.63110 (12)	0.50566 (11)	0.0548 (4)	

C12	0.4042 (3)	0.62977 (12)	0.44420 (11)	0.0560 (4)
C13	0.3893 (3)	0.67568 (12)	0.36151 (11)	0.0548 (4)
C14	0.5200 (2)	0.67695 (10)	0.29619 (10)	0.0486 (4)
C15	0.4674 (2)	0.69629 (11)	0.20060 (10)	0.0562 (4)
C16	0.5861 (2)	0.69706 (11)	0.13708 (11)	0.0546 (4)
C17	0.75962 (19)	0.67906 (10)	0.16958 (10)	0.0494 (4)
C18	0.8178 (2)	0.66060 (11)	0.26364 (11)	0.0571 (5)
C19	0.6975 (2)	0.65988 (11)	0.32628 (11)	0.0562 (4)
H1	-0.0568	0.5784	0.5357	0.0685*
H2	-0.2733	0.5725	0.6333	0.0775*
H3	-0.2005	0.5738	0.7939	0.0780*
Н5	0.0277	0.5755	0.9364	0.0787*
H6	0.3146	0.5749	1.0017	0.0850*
H7	0.5346	0.5754	0.9055	0.0801*
H8	0.4655	0.5779	0.7442	0.0679*
H15	0.3498	0.7089	0.1793	0.0674*
H16	0.5497	0.7095	0.0732	0.0655*
H18	0.9359	0.6489	0.2843	0.0686*
H19	0.7353	0.6478	0.3901	0.0674*
H13	0.284 (3)	0.7118 (12)	0.3417 (11)	0.064 (5)*
H12	0.507 (3)	0.5928 (12)	0.4638 (12)	0.072 (6)*
H11	0.169 (3)	0.6735 (11)	0.4877 (11)	0.062 (5)*

Atomic displacement parameters $(Å^2)$

	U^{11}	U^{22}	U^{33}	U^{12}	U^{13}	U^{23}
01	0.0896 (10)	0.1221 (13)	0.0611 (8)	0.0165 (8)	0.0280 (8)	0.0112 (8)
O2	0.0506 (8)	0.1242 (12)	0.1018 (11)	0.0034 (8)	0.0215 (8)	0.0112 (9)
N1	0.0591 (8)	0.0612 (9)	0.0492 (8)	0.0021 (7)	0.0141 (7)	-0.0007 (7)
N2	0.0625 (10)	0.0652 (10)	0.0666 (10)	-0.0003 (8)	0.0182 (8)	0.0005 (8)
C1	0.0591 (10)	0.0585 (10)	0.0523 (9)	0.0027 (8)	0.0032 (8)	-0.0049 (8)
C2	0.0506 (10)	0.0699 (12)	0.0729 (12)	0.0021 (9)	0.0070 (9)	-0.0074 (9)
C3	0.0546 (10)	0.0716 (12)	0.0732 (12)	-0.0002 (9)	0.0235 (9)	-0.0052 (9)
C4	0.0587 (10)	0.0492 (9)	0.0541 (10)	0.0008 (7)	0.0171 (8)	-0.0025 (7)
C5	0.0788 (13)	0.0671 (11)	0.0552 (10)	-0.0008 (10)	0.0247 (10)	-0.0015 (9)
C6	0.0938 (15)	0.0728 (12)	0.0456 (9)	-0.0023 (11)	0.0087 (10)	0.0031 (8)
C7	0.0653 (11)	0.0727 (12)	0.0591 (10)	-0.0018 (9)	-0.0020 (9)	0.0068 (9)
C8	0.0553 (10)	0.0576 (10)	0.0572 (10)	0.0014 (8)	0.0094 (8)	0.0050 (8)
C9	0.0510 (9)	0.0427 (8)	0.0506 (9)	0.0012 (7)	0.0099 (8)	0.0001 (7)
C10	0.0545 (9)	0.0465 (9)	0.0483 (9)	0.0029 (7)	0.0116 (8)	-0.0019 (7)
C11	0.0607 (10)	0.0556 (10)	0.0487 (9)	0.0012 (9)	0.0103 (8)	-0.0024 (8)
C12	0.0608 (11)	0.0573 (10)	0.0510 (10)	0.0006 (9)	0.0116 (9)	-0.0004 (8)
C13	0.0578 (10)	0.0553 (10)	0.0518 (10)	0.0021 (8)	0.0096 (8)	0.0003 (8)
C14	0.0532 (9)	0.0456 (9)	0.0469 (9)	0.0002 (7)	0.0064 (7)	0.0010 (7)
C15	0.0486 (9)	0.0686 (11)	0.0504 (9)	0.0068 (8)	0.0040 (8)	0.0046 (8)
C16	0.0551 (10)	0.0633 (10)	0.0444 (9)	0.0030 (8)	0.0039 (8)	0.0037 (8)
C17	0.0494 (9)	0.0478 (9)	0.0525 (9)	-0.0008 (7)	0.0121 (8)	0.0011 (7)
C18	0.0479 (9)	0.0620 (11)	0.0594 (10)	0.0011 (8)	0.0003 (8)	0.0054 (8)

C19	0.0580 (10)	0.0645 (11)	0.0437 (9)	0.0002 (8)	-0.0014 (8)	0.0050 (8)	
Geomet	Geometric parameters (Å, °)						
01—N2	2	1.224 (2)		C14—C19		1.395 (3)	
O2—N2	2	1.222 (2)		C15—C16		1.375 (3)	
N1—C1	10	1.414 (3)		C16—C17		1.374 (2)	
N1—C1	11	1.277 (2)		C17—C18		1.377 (2)	
N2—C1	17	1.464 (3)		C18—C19		1.374 (3)	
C1C2	2	1.400 (3)		C1—H1		0.930	
C1—C1	10	1.375 (2)		С2—Н2		0.930	
C2—C3	3	1.358 (3)		С3—Н3		0.930	
C3—C4	1	1.412 (3)		С5—Н5		0.930	
C4—C5	5	1.416 (3)		С6—Н6		0.930	
C4—C9)	1.417 (3)		С7—Н7		0.930	
C5—Cé	5	1.355 (3)		C8—H8		0.930	
C6—C7	7	1.400 (3)		C11—H11		0.989 (16)	
С7—С8	3	1.364 (3)		C12—H12		0.953 (17)	
C8—C9)	1.416 (2)		C13—H13		0.959 (17)	
C9—C1	10	1.435 (2)		C15—H15		0.930	
С11—С	C12	1.438 (3)		C16—H16		0.930	
C12—C	C13	1.330 (3)		C18—H18		0.930	
C13—C	C14	1.464 (3)		C19—H19		0.930	
C14—C	215	1.392 (2)					
01…C1	6	2.708 (3)		C7···H13 ^{viii}		3.134 (17)	
01…C1	8	3.547 (2)		C8…H3 ^v		3.2394	
O2…C1	6	3.540 (2)		C8…H15 ^{viii}		3.2947	
O2…C1	8	2.729 (3)		C8…H13 ^{viii}		3.148 (17)	
N1…C8		2.840 (2)		C9…H15 ^{viii}		3.3151	
N1…C1	3	3.591 (3)		C9…H18 ^{iv}		3.3321	
C1…C4		2.789 (3)		C9…H13 ^{viii}		3.272 (16)	
C1…C1	1	2.910 (3)		C10····H1 ^{xi}		3.3316	
C2…C9		2.808 (2)		C10···H15 ^{viii}		3.2996	
C3…C1	0	2.798 (3)		C10···H18 ^{iv}		3.4799	
C4…C7		2.797 (3)		C10H19 ^{iv}		3.3466	
C5…C8		2.775 (3)		C11…H1 ^{xi}		3.3541	
C6…C9		2.795 (3)		C11····H2 ^{xi}		3.4512	
C9…C1	1	3.523 (3)		C11…H15 ^{viiii}		3.3164	
C12…C	19	3.029 (3)		C11…H16 ^{viii}		3.1354	
C14…C	17	2.758 (3)		$C11 \cdots H12^{iv}$		3.540 (17)	
C15…C	18	2.767 (3)		$C12 \cdots H2^{v}$		3.4901	
C16…C	19	2.761 (3)		C12····H2 ^{xi}		3.1301	
01…C1	i	3.426 (3)		C12····H16 ^{viii}		3.0046	
01…C5	ii	3.422 (3)		$C12\cdots H12^{iv}$		3.386 (17)	
01…C1	9 ⁱⁱⁱ	3.412 (2)		C13····H2 ^{xi}		3.5604	
O2…C5	iv	3.559 (3)		C13····H16 ^{viii}		3.4816	
02…C1	5 ^v	3.326 (2)		C13…H18 ^{ix}		3.5298	

supporting information

N1…C19iv $3.591 (2)$ C14…H8ivN2…C1i $3.405 (2)$ C14…H8iiiN2…C2i $3.509 (3)$ C15…H6xN2…C5iv $3.566 (3)$ C15…H8iiiC1…O1vi $3.426 (3)$ C16…H2iC1…N2vi $3.405 (2)$ C16…H6xC2…N2vi $3.509 (3)$ C17…H2iC2…C17vi $3.480 (3)$ C19…H1vC4…C18iv $3.559 (3)$ H1…N1xiC5…O1vii $3.422 (3)$ H1…N1xiC5…O2iv $3.566 (3)$ H1…C1xiC7…C13viii $3.456 (3)$ H1…C10xiC8…C14viii $3.575 (2)$ H1…C11xiC8…C15viii $3.448 (3)$ H1…C19ixC9…C18iv $3.329 (2)$ H1…H1xi	3.5866 3.4924 3.3647 3.1933 3.3773 3.1359 3.4883 3.4892 3.2020 3.1668 3.5116 3.0465
N2···C1i $3.405(2)$ $C14H8^{iii}$ N2···C2i $3.509(3)$ $C15H6^x$ N2···C5iv $3.566(3)$ $C15H8^{iii}$ C1···01^{vi} $3.426(3)$ $C16H2^i$ C1···N2^{vi} $3.405(2)$ $C16H6^x$ C2···N2^{vi} $3.509(3)$ $C17H2^i$ C2···C17^{vi} $3.480(3)$ $C19H1^v$ C4···C18^{iv} $3.561(3)$ $H1N1^{xi}$ C5···O1^{vii} $3.422(3)$ $H1N2^{vi}$ C5···O2^{iv} $3.559(3)$ $H1N2^{vi}$ C5···N2^{iv} $3.566(3)$ $H1C1^{xi}$ C7···C13^{viii} $3.456(3)$ $H1C10^{si}$ C8···C14^{viii} $3.575(2)$ $H1C19^{ix}$ C9···C18^{iv} $3.329(2)$ $H1H1^{xi}$	3.4924 3.3647 3.1933 3.3773 3.1359 3.4883 3.4892 3.2020 3.1668 3.5116 3.0465
N2…C2i $3.509(3)$ C15…H6xN2…C5iv $3.566(3)$ C15…H8iiiC1…O1vi $3.426(3)$ C16…H2iC1…N2vi $3.405(2)$ C16…H6xC2…N2vi $3.509(3)$ C17…H2iC2…C17vi $3.480(3)$ C19…H1vC4…C18iv $3.561(3)$ H1…O1viC5…O1vii $3.422(3)$ H1…N1xiC5…O2iv $3.559(3)$ H1…N2viC5…N2iv $3.566(3)$ H1…C1xiC7…C13viii $3.456(3)$ H1…C10xiC8…C14viii $3.575(2)$ H1…C11xiC8…C15viii $3.448(3)$ H1…C19ixC9…C18iv $3.329(2)$ H1…H1xi	3.3647 3.1933 3.3773 3.1359 3.4883 3.4892 3.2020 3.1668 3.5116 3.0465
N2···C5 ^{iv} 3.566 (3) $C15 \cdots H8^{iii}$ C1···O1 ^{vi} 3.426 (3) $C16 \cdots H2^i$ C1···N2 ^{vi} 3.405 (2) $C16 \cdots H6^s$ C2···N2 ^{vi} 3.509 (3) $C17 \cdots H2^i$ C2···C17 ^{vi} 3.480 (3) $C19 \cdots H1^v$ C4···C18 ^{iv} 3.561 (3) $H1 \cdots O1^{vi}$ C5···O1 ^{vii} 3.422 (3) $H1 \cdots N1^{xi}$ C5···O2 ^{iv} 3.559 (3) $H1 \cdots N2^{vi}$ C5···N2 ^{iv} 3.566 (3) $H1 \cdots C1^{xi}$ C7···C13 ^{viii} 3.456 (3) $H1 \cdots C10^{xi}$ C8···C14 ^{viii} 3.575 (2) $H1 \cdots C19^{ix}$ C9···C18 ^{iv} 3.329 (2) $H1 \cdots H1^{xi}$	3.1933 3.3773 3.1359 3.4883 3.4892 3.2020 3.1668 3.5116 3.0465
$C1\cdots O1^{vi}$ 3.426 (3) $C16\cdots H2^{i}$ $C1\cdots N2^{vi}$ 3.405 (2) $C16\cdots H6^{x}$ $C2\cdots N2^{vi}$ 3.509 (3) $C17\cdots H2^{i}$ $C2\cdots C17^{vi}$ 3.480 (3) $C19\cdots H1^{v}$ $C4\cdots C18^{iv}$ 3.561 (3) $H1\cdots O1^{vi}$ $C5\cdots O1^{vii}$ 3.422 (3) $H1\cdots N1^{xi}$ $C5\cdots O2^{iv}$ 3.559 (3) $H1\cdots N2^{vi}$ $C5\cdots N2^{iv}$ 3.566 (3) $H1\cdots C1^{xi}$ $C7\cdots C13^{viii}$ 3.456 (3) $H1\cdots C10^{xi}$ $C8\cdots C14^{viii}$ 3.575 (2) $H1\cdots C11^{xi}$ $C8\cdots C15^{viii}$ 3.448 (3) $H1\cdots C19^{ix}$ $C9\cdots C18^{iv}$ 3.329 (2) $H1\cdots H1^{xi}$	3.3773 3.1359 3.4883 3.4892 3.2020 3.1668 3.5116 3.0465
$C1 \cdots N2^{vi}$ $3.405 (2)$ $C16 \cdots H6^{x}$ $C2 \cdots N2^{vi}$ $3.509 (3)$ $C17 \cdots H2^{i}$ $C2 \cdots C17^{vi}$ $3.480 (3)$ $C19 \cdots H1^{v}$ $C4 \cdots C18^{iv}$ $3.561 (3)$ $H1 \cdots O1^{vi}$ $C5 \cdots O1^{vii}$ $3.422 (3)$ $H1 \cdots N1^{xi}$ $C5 \cdots O2^{iv}$ $3.559 (3)$ $H1 \cdots N2^{vi}$ $C5 \cdots N2^{iv}$ $3.566 (3)$ $H1 \cdots C1^{xi}$ $C7 \cdots C13^{viii}$ $3.456 (3)$ $H1 \cdots C10^{xi}$ $C8 \cdots C14^{viii}$ $3.575 (2)$ $H1 \cdots C11^{xi}$ $C8 \cdots C15^{viii}$ $3.448 (3)$ $H1 \cdots C19^{ix}$ $C9 \cdots C18^{iv}$ $3.329 (2)$ $H1 \cdots H1^{xi}$	3.1359 3.4883 3.4892 3.2020 3.1668 3.5116 3.0465
$C2\cdots N2^{vi}$ $3.509(3)$ $C17\cdots H2^{i}$ $C2\cdots C17^{vi}$ $3.480(3)$ $C19\cdots H1^{v}$ $C4\cdots C18^{iv}$ $3.561(3)$ $H1\cdots O1^{vi}$ $C5\cdots O1^{vii}$ $3.422(3)$ $H1\cdots N1^{xi}$ $C5\cdots O2^{iv}$ $3.559(3)$ $H1\cdots N2^{vi}$ $C5\cdots N2^{iv}$ $3.566(3)$ $H1\cdots C1^{xi}$ $C7\cdots C13^{viii}$ $3.456(3)$ $H1\cdots C10^{xi}$ $C8\cdots C14^{viii}$ $3.575(2)$ $H1\cdots C11^{xi}$ $C8\cdots C15^{viii}$ $3.448(3)$ $H1\cdots C19^{ix}$ $C9\cdots C18^{iv}$ $3.329(2)$ $H1\cdots H1^{xi}$	3.4883 3.4892 3.2020 3.1668 3.5116 3.0465
$C2\cdots C17^{vi}$ 3.480 (3) $C19\cdots H1^v$ $C4\cdots C18^{iv}$ 3.561 (3) $H1\cdots O1^{vi}$ $C5\cdots O1^{vii}$ 3.422 (3) $H1\cdots N1^{xi}$ $C5\cdots O2^{iv}$ 3.559 (3) $H1\cdots N2^{vi}$ $C5\cdots N2^{iv}$ 3.566 (3) $H1\cdots C1^{xi}$ $C7\cdots C13^{viii}$ 3.456 (3) $H1\cdots C1^{xi}$ $C8\cdots C14^{viii}$ 3.575 (2) $H1\cdots C11^{xi}$ $C8\cdots C15^{viii}$ 3.448 (3) $H1\cdots C19^{ix}$ $C9\cdots C18^{iv}$ 3.329 (2) $H1\cdots H1^{xi}$	3.4892 3.2020 3.1668 3.5116 3.0465
$C4\cdots C18^{iv}$ $3.561 (3)$ $H1\cdots O1^{vi}$ $C5\cdots O1^{vii}$ $3.422 (3)$ $H1\cdots N1^{xi}$ $C5\cdots O2^{iv}$ $3.559 (3)$ $H1\cdots N2^{vi}$ $C5\cdots N2^{iv}$ $3.566 (3)$ $H1\cdots C1^{xi}$ $C7\cdots C13^{viii}$ $3.456 (3)$ $H1\cdots C10^{xi}$ $C8\cdots C14^{viii}$ $3.575 (2)$ $H1\cdots C11^{xi}$ $C8\cdots C15^{viii}$ $3.448 (3)$ $H1\cdots C19^{ix}$ $C9\cdots C18^{iv}$ $3.329 (2)$ $H1\cdots H1^{xi}$	3.2020 3.1668 3.5116 3.0465
$C5\cdots O1^{vii}$ $3.422 (3)$ $H1\cdots N1^{xi}$ $C5\cdots O2^{iv}$ $3.559 (3)$ $H1\cdots N2^{vi}$ $C5\cdots N2^{iv}$ $3.566 (3)$ $H1\cdots C1^{xi}$ $C7\cdots C13^{viii}$ $3.456 (3)$ $H1\cdots C10^{xi}$ $C8\cdots C14^{viii}$ $3.575 (2)$ $H1\cdots C11^{xi}$ $C8\cdots C15^{viii}$ $3.448 (3)$ $H1\cdots C19^{ix}$ $C9\cdots C18^{iv}$ $3.329 (2)$ $H1\cdots H1^{xi}$	3.1668 3.5116 3.0465
$C5\cdots O2^{iv}$ $3.559 (3)$ $H1\cdots N2^{vi}$ $C5\cdots N2^{iv}$ $3.566 (3)$ $H1\cdots C1^{xi}$ $C7\cdots C13^{viii}$ $3.456 (3)$ $H1\cdots C10^{xi}$ $C8\cdots C14^{viii}$ $3.575 (2)$ $H1\cdots C11^{xi}$ $C8\cdots C15^{viii}$ $3.448 (3)$ $H1\cdots C19^{ix}$ $C9\cdots C18^{iv}$ $3.329 (2)$ $H1\cdots H1^{xi}$	3.5116 3.0465
$C5\cdots N2^{iv}$ $3.566 (3)$ $H1\cdots C1^{xi}$ $C7\cdots C13^{viii}$ $3.456 (3)$ $H1\cdots C10^{xi}$ $C8\cdots C14^{viii}$ $3.575 (2)$ $H1\cdots C11^{xi}$ $C8\cdots C15^{viii}$ $3.448 (3)$ $H1\cdots C19^{ix}$ $C9\cdots C18^{iv}$ $3.329 (2)$ $H1\cdots H1^{xi}$	3.0465
$C7\cdots C13^{viii}$ 3.456 (3) $H1\cdots C10^{xi}$ $C8\cdots C14^{viii}$ 3.575 (2) $H1\cdots C11^{xi}$ $C8\cdots C15^{viii}$ 3.448 (3) $H1\cdots C19^{ix}$ $C9\cdots C18^{iv}$ 3.329 (2) $H1\cdots H1^{xi}$	
$C8 \cdots C14^{viii}$ $3.575 (2)$ $H1 \cdots C11^{xi}$ $C8 \cdots C15^{viii}$ $3.448 (3)$ $H1 \cdots C19^{ix}$ $C9 \cdots C18^{iv}$ $3.329 (2)$ $H1 \cdots H1^{xi}$	3.3316
$C8 \cdots C15^{viii}$ $3.448 (3)$ $H1 \cdots C19^{ix}$ $C9 \cdots C18^{iv}$ $3.329 (2)$ $H1 \cdots H1^{xi}$	3.3541
C9…C18 ^{iv} 3.329 (2) H1…H1 ^{xi}	3.4892
	2.6052
C9C19 ^{iv} $3.564(3)$ H1H19 ^{ix}	2.6192
$C10C19^{iv}$ 3.567 (2) $H1H12^{ix}$	3.3760
C13C7 ⁱⁱⁱ 3.456 (3) $H2N1^{ix}$	3.3669
C14C8 ⁱⁱⁱ $3.575(2)$ H2C11 ^{xi}	3.4512
$C15O2^{ix}$ 3.326 (2) $H2C12^{ix}$	3.4901
C15C8 ⁱⁱⁱ $3.448(3)$ H2C12 ^{xi}	3.1301
C17C2 ⁱ $3.480(3)$ H2C13 ^{xi}	3.5604
C18···C4 ^{iv} $3.561(3)$ H2···C16 ^{vi}	3.3773
C18···C9 ^{iv} $3.329(2)$ H2···C17 ^{vi}	3.4883
C19O1 ^{viii} $3.412(2)$ $H2H8^{ix}$	2.7294
C19N1 ^{iv} $3.591(2)$ H2H16 ^{vi}	3.3765
C19C9 ^{iv} $3.564(3)$ H2H12 ^{ix}	2.7617
C19C10 ^{iv} $3.567(2)$ H2H12 ^{xi}	3.1167
Q1H16 2.4154 H3C7 ^{ix}	3.3298
O2…H18 2.4477 H3…C8 ^{ix}	3.2394
N1…H1 2.6358 H3…H7 ^{ix}	2.7580
N1···H8 2.5219 H3···H8 ^{ix}	2.5703
N1···H12 2.558 (19) H5···O1 ^{vii}	2.6614
N2…H16 2.5955 H5…O2 ^{vii}	3.0276
N2…H18 2.6122 H5…O2 ^{iv}	3.4966
C1···H3 3.2302 H5···N2 ^{vii}	3.1031
C1···H11 2.695 (17) H5···C5 ^{xii}	3.4610
C3···H1 3.2230 H5···H5 ^{xii}	2.8416
C3···H5 2.6512 H5···H6 ^{xii}	3.5690
	3.2431
J_{112} J_{12} J_{12} J_{12}	3.2526
C4···H6 3.2546 H6···C7 ^{xiii}	3.3647
C4···H6 3.25460 H6···C2 C4···H8 3.2801 H6···C15 ^{xiv}	
C4···H2 3.2400 H0···O2 C4···H6 3.2546 H6···C7 ^{xiii} C4···H8 3.2801 H6···C15 ^{xiv} C5···H3 2.6530 H6···C16 ^{xiv}	3.1359
C4···H2 3.2460 H0···O2 C4···H6 3.2546 H6···C7 ^{xiii} C4···H8 3.2801 H6···C15 ^{xiv} C5···H3 2.6530 H6···C16 ^{xiv} C5···H7 3.2228 H6···H5 ^{xii}	3.1359 3.5690
C4···H2 3.2460 H0···O2 C4···H6 3.2546 H6···C7 ^{xiii} C4···H8 3.2801 H6···C15 ^{xiv} C5···H3 2.6530 H6···C16 ^{xiv} C5···H7 3.2228 H6···H5 ^{xii} C6···H8 3.2377 H6···H6 ^{xiii}	3.1359 3.5690 3.5374
C4 ···H2 3.2460 H0···O2 C4···H6 3.2546 H6···C7 ^{xiii} C4···H8 3.2801 H6···C15 ^{xiv} C5···H3 2.6530 H6···C16 ^{xiv} C5···H7 3.2228 H6···H5 ^{xii} C6···H8 3.2377 H6···H6 ^{xiii} C7···H5 3.2269 H6···H7 ^{xiii}	3.1359 3.5690 3.5374 2.6476

С9…Н1	3.2677	H6…H16 ^{xiv}	2.6995
С9…Н3	3.2819	H7…O1 ^{xiv}	3.1257
С9…Н5	3.2729	H7····C6 ^{xiii}	3.2438
С9…Н7	3.2542	H7····H3 ^v	2.7580
С10…Н2	3.2503	H7…H6 ^{xiii}	2.6476
С10…Н8	2.6791	H7…H7 ^{xiii}	3.5092
C10…H11	2.515 (16)	H7···H16 ^{xiv}	3.0192
С11…Н1	2 7219	H7···H13 ^{viii}	3 5723
C11····H8	3 5939	H8····C2 ^v	3 3067
C11H13	2 607 (16)	H8···C3 ^v	3 2321
C12H19	2.007 (10)	H8···C14 ^{iv}	3 5866
C12H15	2.1179	H8···C14 ^{viii}	3 4924
C13H10	2.0129		3.1023
C13H11	2.0070 2.642(17)	По ^{те} ст5 Ц8Ц2v	2 7204
	2.042(17)		2.7234
C14H18	3.2301		2.3703
C14H18	3.2332		3.18/3
C14H12	2.6/3 (1/)		3.5949
С15…Н19	3.2306		2.4604
С15…Н13	2.622 (17)		3.2294
С16…Н18	3.2430	H15C8 ^m	3.2947
С17…Н15	3.2058	H15…C9 ^m	3.3151
С17…Н19	3.2044	H15…C10 ^m	3.2996
C18…H16	3.2425	H15…C11 ⁱⁱⁱ	3.3164
С19…Н15	3.2315	H15…H6 ^x	3.1225
С19…Н13	3.300 (18)	H15…H8 ⁱⁱⁱ	3.1875
С19…Н12	2.769 (18)	H15…H11 ⁱⁱⁱ	3.3154
H1…H2	2.3187	H16…N1 ⁱⁱⁱ	3.5481
H1…H11	2.3587	H16···C6 ^x	3.2099
Н2…Н3	2.2795	H16····C7 ^x	3.3756
Н3…Н5	2.4921	H16…C11 ⁱⁱⁱ	3.1354
Н5…Н6	2.2740	H16…C12 ⁱⁱⁱ	3.0046
H6…H7	2.3246	H16…C13 ⁱⁱⁱ	3.4816
H7…H8	2.2861	H16…H2 ⁱ	3.3765
H15…H16	2.3051	H16…H6 ^x	2.6995
H15…H13	2.4371	H16…H7 ^x	3.0192
H18…H19	2.3055	H16…H12 ⁱⁱⁱ	3.1495
H19…H13	3.5569	H16…H11 ⁱⁱⁱ	3.4229
H19…H12	2.3008	H18····C1 ^{iv}	3.5734
H13···H12	2.80(3)	H18····C2 ^{iv}	3 5731
H13···H11	2.00(3) 2 44 (3)	H18····C3 ^{iv}	3 4701
H12···H11	2.00(3)	H18····C4 ^{iv}	3 3461
01···H1 ⁱ	3 2020	H18Coiv	3 3321
01H5 ⁱⁱ	2 6614	H18····C10 ^{iv}	3 4799
$01 \cdots H7^{x}$	3 1257	H18C13 ^v	3 5208
$01 \dots H10^{iii}$	2 8 3 8 7	H18H12v	2.2270 2.8215
$01 \dots 1117$	2.0307	ш18Ш11v	2.0313
$O_1 IIII$	3.217(17)		2.2024 2.2024
$O_2 II_3 O_2 II_3 O_2 II_3 O_2 II_3 O_2 II_3 O_2 O_2 O_2 O_2 O_2 O_2 O_2 O_2 O_2 O_2$	3.0270		2.030/
04 113	J.+700	1117 TINI	J.414/

02…H6 ⁱⁱ	3 2431	H19C1 ^v	3 4374
$02 \cdot H0$	2 4604	H19····C10 ^{iv}	3 3466
02° H13	3 375 (15)	H19H1v	2 6192
$02H11^{i}$	3 269 (16)	H19H11 ^v	3 4503
N1H1 ^{xi}	3 1668	$H13O2^{ix}$	3,375 (15)
N1H2v	3 3660	H13C/ ⁱⁱⁱ	3.375(17)
N1H15 ^{viii}	3 2294	H13C5 ⁱⁱⁱ	3.394(17)
N1H16 ^{viii}	3 5/81	H13C6 ⁱⁱⁱ	3.300(17)
	3.3401	H13 C0	3.241(10) 3.134(17)
N1H12iv	3.2127	H13·C8iii	3.134(17)
N2H1i	3 5116		3.140(17) 3.272(16)
N2U5 ⁱⁱ	2 1021	H13 C9 H13H7 ⁱⁱⁱ	3.272 (10)
N2U11 ⁱ	3,530 (17)		3.5725
112^{-11111}	3.0465	П15 116 Ц12Ц19 ^{ix}	2 8215
	2 5724	1115 1118	2.0313
	2 4274		3.013(17)
	2.20(7		5.545 (10) 2.540 (17)
$C_2 = H_1 e_{ix}$	3.300/		3.340(17)
C2H18 ⁱⁿ	5.5/51 2.545 (1()		3.386 (17)
C2. Usix	5.545 (10) 2.2221		3.3760
	3.2321		2.7617
	3.4701		3.1167
	3.3461		3.1495
	3.394 (17)		2.78 (3)
	3.4610		3.219 (17)
C5H13vm	3.360 (17)		3.269 (16)
C6···H7 ^{xm}	3.2438	H11N2 ^{v1}	3.530 (17)
C6…H16 ^{xiv}	3.2099	H11····H15 ^{viii}	3.3154
C6···H13 ^{viii}	3.241 (16)	H11···H16 ^{viii}	3.4229
С7…Н3 ^v	3.3298	H11···H18 ^{ix}	3.2054
C7···H6 ^{xiii}	3.2526	H11···H19 ^{ix}	3.4503
C7···H16 ^{xiv}	3.3756		
C10—N1—C11	119.53 (14)	C17—C18—C19	118.51 (15)
O1—N2—O2	122.81 (17)	C14—C19—C18	121.37 (14)
01—N2—C17	118.37 (14)	C2-C1-H1	119.358
O2—N2—C17	118.82 (15)	С10—С1—Н1	119.366
C2-C1-C10	121.28 (15)	С1—С2—Н2	119.856
C1—C2—C3	120.30 (15)	C3—C2—H2	119.847
C2—C3—C4	120.87 (17)	С2—С3—Н3	119.562
C3—C4—C5	122.11 (17)	C4—C3—H3	119.566
C3—C4—C9	119.53 (15)	C4—C5—H5	119.342
C5-C4-C9	118.36 (15)	С6—С5—Н5	119.342
C4—C5—C6	121.32 (18)	C5—C6—H6	119.866
C5—C6—C7	120.27 (16)	C7—C6—H6	119.866
C6-C7-C8	120.49 (16)	C6—C7—H7	119.748
C7-C8-C9	120.61 (16)	C8—C7—H7	119 758
C4-C9-C8	118 94 (14)	C7—C8—H8	119 691
C4-C9-C10	118 51 (13)	C9 - C8 - H8	119 700
	110.01 (10)	07 00 110	117.700

122.54 (15)	N1—C11—H11	121.1 (10)
123.40 (13)	C12—C11—H11	117.4 (10)
116.99 (13)	C11—C12—H12	117.6 (11)
119.47 (15)	C13—C12—H12	118.8 (11)
121.48 (16)	С12—С13—Н13	118.4 (10)
123.62 (16)	C14—C13—H13	115.6 (10)
126.01 (16)	C14—C15—H15	119.424
119.57 (14)	C16—C15—H15	119.428
122.30 (14)	С15—С16—Н16	120.586
118.13 (15)	C17—C16—H16	120.584
121.15 (14)	C17—C18—H18	120.741
118.83 (14)	C19—C18—H18	120.752
118.69 (13)	C14—C19—H19	119.312
119.29 (14)	C18—C19—H19	119.319
122.01 (15)		
-177.42 (12)	C6—C7—C8—C9	0.2 (3)
41.0 (2)	C7—C8—C9—C4	-1.1 (2)
-143.39 (13)	C7—C8—C9—C10	179.35 (13)
0.2 (2)	C4—C9—C10—N1	-178.53 (11)
-178.89 (13)	C4—C9—C10—C1	-2.72 (19)
179.70 (13)	C8—C9—C10—N1	1.1 (2)
0.6 (2)	C8—C9—C10—C1	176.87 (12)
177.29 (13)	N1-C11-C12-C13	175.92 (14)
1.8 (3)	C11—C12—C13—C14	179.95 (14)
0.1 (3)	C12—C13—C14—C15	156.74 (15)
-1.0 (3)	C12—C13—C14—C19	-23.6 (3)
-179.08 (14)	C13—C14—C15—C16	-179.30 (13)
-0.0 (3)	C13—C14—C19—C18	179.44 (13)
178.28 (14)	C15—C14—C19—C18	-0.9 (2)
-177.73 (12)	C19—C14—C15—C16	1.1 (2)
1.9 (2)	C14—C15—C16—C17	-0.5 (3)
1.4 (2)	C15—C16—C17—N2	-179.27 (12)
-179.04 (12)	C15—C16—C17—C18	-0.2 (3)
-0.8 (3)	N2-C17-C18-C19	179.39 (12)
-0.1 (3)	C16—C17—C18—C19	0.3 (3)
0.4 (3)	C17—C18—C19—C14	0.3 (3)
	122.54 (15) $123.40 (13)$ $116.99 (13)$ $119.47 (15)$ $121.48 (16)$ $123.62 (16)$ $126.01 (16)$ $119.57 (14)$ $122.30 (14)$ $118.13 (15)$ $121.15 (14)$ $118.83 (14)$ $118.69 (13)$ $119.29 (14)$ $122.01 (15)$ $-177.42 (12)$ $41.0 (2)$ $-143.39 (13)$ $0.2 (2)$ $-178.89 (13)$ $179.70 (13)$ $0.6 (2)$ $177.29 (13)$ $1.8 (3)$ $0.1 (3)$ $-179.08 (14)$ $-0.0 (3)$ $178.28 (14)$ $-177.73 (12)$ $1.9 (2)$ $1.4 (2)$ $-0.1 (3)$ $0.4 (3)$	122.54 (15) N1—C11—H11 $123.40 (13)$ C12—C11—H11 $116.99 (13)$ C11—C12—H12 $119.47 (15)$ C13—C12—H12 $119.47 (15)$ C13—C12—H12 $121.48 (16)$ C12—C13—H13 $123.62 (16)$ C14—C15—H15 $119.57 (14)$ C16—C15—H15 $122.30 (14)$ C15—C16—H16 $118.13 (15)$ C17—C16—H16 $121.15 (14)$ C17—C18—H18 $118.69 (13)$ C14—C19—H19 $122.01 (15)$ C6—C7—C8—C9 $-177.42 (12)$ C6—C7—C8—C9—C10 $0.2 (2)$ C4—C9—C10—N1 $-178.89 (13)$ C4—C9—C10—N1 $-178.89 (13)$ C4—C9—C10—N1 $-177.29 (13)$ N1—C11—C12—C13 $1.8 (3)$ C11—C12—C13—C14 $0.1 (3)$ C12—C13—C14—C15 $-1.0 (3)$ C12—C13—C14—C19 $-177.73 (12)$ C19—C14—C15—C16 $-179.04 (12)$ C15—C16—C17—N2 $-179.04 (12)$ C15—C16—C17—C18 $-18.(3)$ N2—C17—C18—C19 $-0.1 (3)$ C16—C17—C18—C19 $-0.1 (3)$ C16—C17—C18—C1

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

Hydrogen-bond geometry (Å, °)

D—H···A	D—H	H···A	$D \cdots A$	D—H···A
C5—H5····O1 ^{vii}	0.93	2.66	3.422 (3)	139
C15—H15…O2 ^{ix}	0.93	2.46	3.326 (3)	155

Symmetry codes: (vii) x-1, y, z+1; (ix) x-1, y, z.