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# Crystal structure of (E)-2-hydroxy-1,2-diphenyl-ethan-1-one oxime 

Hans Reuter ${ }^{\text {a* }}$ and Coco K. Y. A. Okio ${ }^{\text {b }}$

${ }^{\text {a }}$ Institute of Chemistry of New Materials, University of Osnabrück, Barbarastrasse 7, 49069 Osnabrück, Germany, and ${ }^{\text {b }}$ Departamento de Química, Facultad de Ciencias, Universidad Nacional de Colombia, Carerra 30 No 45-03, Bogotá, Colombia. *Correspondence e-mail: hreuter@uos.de

The title compound, $\mathrm{C}_{14} \mathrm{H}_{13} \mathrm{NO}_{2}$, is a commercially available material and can be used as a multidentate ligand. The molecule of the asymmetric unit has an $R$ configuration, while the corresponding $S$-configured molecule of the racemic mixture is generated by a crystallographic centre of symmetry. Both hydroxy groups (the H atom of the oxime group is equally disordered over two positions) are involved in hydrogen bonding, leading to the formation of chains extending parallel to [001].

## 1. Chemical context

The title compound ( $E$ )-2-hydroxy-1,2-diphenyl-ethan-1-one oxime, $\mathrm{C}_{14} \mathrm{H}_{13} \mathrm{NO}_{2}$, is commercially available and can be used as a multidentate ligand for which many trivial names such as cuprone or alpha-benzoin, and abbreviations including $\mathrm{AboH}_{2}, \mathrm{BzoxH}_{2}$, are in use. Used for a long time for the determination of manganese or copper in steel (Feigl, 1923; Knowles, 1932; Kar, 1935), $\mathrm{BzoxH}_{2}$ has attracted considerable attention nowadays in the coordination chemistry of transition metals for the preparation of molecular wheels and highnuclearity metal units with copper, manganese or nickel cations (Stamatatos et al., 2012; Vlahopoulou et al., 2009; Koumousi et al. 2010; Karotsis et al., 2009). In the course of a project to evaluate the reactivity of $\mathrm{BzoxH}_{2}$ towards organotin(IV) compounds, we obtained high-quality single crystals of the title compound which we have used for structure determination by X-ray diffraction.


## 2. Structural commentary

BzoxH2 crystallizes in the centrosymmetric monoclinic space group $C 2 / c$ with eight molecules in the unit cell and one molecule in the asymmetric unit. As the compound possesses an asymmetric carbon atom (C2), the molecule of the asymmetric unit has an $R$-configuration while the corresponding $S$-configured molecule of the racemic mixture is generated by a crystallographic centre of symmetry. Both molecules also

Table 1
Selected geometric parameters ( $\left(\AA,{ }^{\circ}\right)$.

| $\mathrm{O} 1-\mathrm{N} 1$ | $1.404(1)$ | $\mathrm{C} 1-\mathrm{C} 2$ | $1.525(2)$ |
| :--- | :---: | :--- | ---: |
| $\mathrm{C} 1-\mathrm{N} 1$ | $1.278(2)$ | $\mathrm{O} 2-\mathrm{C} 2$ | $1.425(2)$ |
|  |  |  |  |
| $\mathrm{N} 1-\mathrm{C} 1-\mathrm{C} 2$ | $114.3(1)$ | $\mathrm{C} 1-\mathrm{N} 1-\mathrm{O} 1$ | $115.5(1)$ |
| $\mathrm{C} 11-\mathrm{C} 1-\mathrm{C} 2$ | $117.7(1)$ | $\mathrm{O} 2-\mathrm{C} 2-\mathrm{C} 1$ | $110.1(1)$ |

Table 2
Hydrogen-bond geometry ( $\AA{ }^{\circ}{ }^{\circ}$ ).

| $D-\mathrm{H} \cdots A$ | $D-\mathrm{H}$ | $\mathrm{H} \cdots A$ | $D \cdots A$ | $D-\mathrm{H} \cdots A$ |
| :--- | :--- | :--- | :--- | :--- |
| $\mathrm{O}^{2}-\mathrm{H} 1 \cdots \mathrm{~N} 1^{\mathrm{i}}$ | 0.96 | 1.97 | $2.805(2)$ | 144 |
| $\mathrm{O}^{\mathrm{ii}}-\mathrm{H} 3 \cdots \mathrm{O} 2^{\text {ii }}$ | 0.96 | 1.89 | $2.829(2)$ | 164 |
| $\mathrm{O}^{\mathrm{O}}-\mathrm{H} 4 \cdots \mathrm{O}^{\mathrm{iii}}$ | 0.96 | 1.85 | $2.806(2)$ | 175 |

Symmetry codes: (i) $-x+1, y,-z+\frac{3}{2}$; (ii) $-x+1,-y,-z+1$; (iii) $-x+1, y,-z+\frac{1}{2}$.
show the $E$ configuration at the $\mathrm{N}=\mathrm{C}$ double bond of the oxime moiety (Fig. 1).

The length [1.278 (2) $\AA$ ] of the $\mathrm{N}=\mathrm{C}$ double bond (Table 1) is consistent with the value of 1.281 (13) $\AA$ found in other oxime moieties (Allen et al., 1987). In addition, this moiety is characterized by a bond angle of $115.5(1)^{\circ}$ at the N atom and of $102.1^{\circ}$ at the O atom. The central $\mathrm{C}-\mathrm{C}$ bond of the molecule has a length of 1.525 (2), which is also in good accordance with a typical single bond between $s p^{3}(\mathrm{C} 2)$ and $s p^{2}(\mathrm{C} 1)$ hybridized C atoms. As a consequence of the different hybridization states, however, the bonds of these two carbon atoms to their phenyl groups are slightly different: 1.512 (2) $\AA$ for C 2 and 1.484 (2) $\AA$ for C1, respectively. The hydroxy group attached to C 2 shows a $\mathrm{C}-\mathrm{O}$ bond length of 1.425 (2) $\AA$, which also lies in the normal range $\left(1.421-1.433 \AA\right.$ ) of a $\mathrm{C}_{2}{ }^{-}$ $\mathrm{CH}-\mathrm{OH}$ group (Allen et al., 1987).
The two phenyl groups exhibit a mean $\mathrm{C}-\mathrm{C}$ bond length of 1.387 (5) $\AA$ [variation: 1.374 (3) -1.398 (2) $\AA$ ], in excellent


Figure 1
The asymmetric unit of the title compound, showing the atom-labelling scheme and displacement ellipsoids for the non-H atoms at the $50 \%$ probability level; split positions of the H atom attached to atom O 2 are labelled H3 and H4.


Figure 2
Crystal packing showing the tube-like arrangement of the molecules along [001].
agreement with the literature value (Allen et al., 1987) of 1.387 (10) $\AA$ for $C_{a r}-C_{a r}$. The mean value of the endocyclic bond angles within the phenyl rings is $120.0(5)^{\circ}$ with minima at the ipso carbon atoms C11 [118.3 (1) $\left.{ }^{\circ}\right]$ and $\mathrm{C} 21\left[119.1(1)^{\circ}\right]$. The phenyl rings form an interplanar angle of $80.72(5)^{\circ}$.

## 3. Supramolecular features

The molecule possesses two hydroxy groups which, in principle, can act as donors and acceptors for hydrogen bonding while the N atom of the oxime moiety can only act as an acceptor atom in the formation of hydrogen bonds. In fact, the crystal packing (Fig. 2) with its clear separation of polar and non-polar moieties, results from two different types of hydrogen bonds (Table 2), giving rise to a one-dimensional tube-like arrangement of the molecules propagating along [001]. In the first type of hydrogen bond, only the hydroxy group attached to the carbon atom C2 is involved, acting both as hydrogen-donor and hydrogen-acceptor groups (Fig. 3). Since the oxygen atoms of the resulting hydrogen bonds are related to each other by a centre of symmetry $\left[\mathrm{O} 2 \cdots \mathrm{O} 2^{\mathrm{ii}}=\right.$


Figure 3
Detail of the one-dimensional hydrogen-bonding system (red dashed lines) derived from the hydroxy group attached to the C atom looking down [010]; displacement ellipsoids for the non-H atoms are drawn at the $50 \%$ probability level. Groups attached to C atoms have been omitted for clarity. Small black dots visualize the position of an inversion center [i1: $\frac{1}{2}, 0,1 ; i 2: \frac{1}{2}, 0, \frac{1}{2} ;$; $\left.3: \frac{1}{2}, 0,0\right]$, green dots the position of twofold rotation axes $\left[\mathrm{r} 1: \frac{1}{2}, y, \frac{3}{4} ; \mathrm{r} 2: \frac{1}{2}, y, \frac{1}{4}\right.$ ]. [Symmetry codes used to generate equivalent atoms: (1) $1-x, y, \frac{1}{2}-z ;$ (2) $x,-y,-\frac{1}{2}+z$; (3) $1-x,-y, 1-z$; (4) $x,-y$, $\frac{1}{2}+z ;$ (5) $1-x, y, \frac{3}{2}-z$.]


Figure 4
Hydrogen-bonding system (red dashed lines) between the oxime groups of two neighbouring molecules looking down [010]; displacement ellipsoids for the non-H atoms are given at the $50 \%$ probability level. The small green dot visualizes the position of the twofold rotation axis at $\frac{1}{2}, y, \frac{3}{4}$. [Symmetry codes used to generate equivalent atoms: (1) $1-x, y$, $\frac{3}{2}-z$.]
$2.829(2) \AA,\left\langle\mathrm{O} 2-\mathrm{H} 3 \cdots \mathrm{O} 2^{\text {ii }}=164^{\circ}\right.$; symmetry code: (ii) $=$ $-x+1,-y,-z+1]$ and a twofold rotation axis $\left[\mathrm{O} 2 \cdots \mathrm{O} 2^{\text {iii }}=\right.$ $2.806(2) \AA,\left\langle\mathrm{O} 2-\mathrm{H} 4 \cdots \mathrm{O} 1^{\mathrm{iii}}=175^{\circ}\right.$; symmetry code (iii) $=$ $\left.-x+1, y,-z+\frac{1}{2}\right]$, respectively, the hydrogen atom of the hydroxy group breaks space-group symmetry, which was considered in the structure model by two equally disordered split positions [ $\mathrm{H} 3 / \mathrm{H} 4]$ of this hydrogen atom. While this kind of hydrogen-bonding system extends to an infinite number of molecules, the second type of hydrogen bond is limited to two neigbouring molecules. It involves the hydroxy group of the oxime moiety that acts as an H -atom donor forming mutual hydrogen bonds with the nitrogen atom of the oxime moiety of a neighbouring molecule, giving rise to two equivalent hydrogen bonds [O1 $\cdots \mathrm{N} 1^{\mathrm{i}}=2.805(2) \AA,\left\langle\mathrm{O} 1-\mathrm{H} 1 \cdots \mathrm{~N} 1^{\mathrm{i}}=\right.$ $144^{\circ}$; symmetry code: (i) $\left.=-x+1, y,-z+\frac{3}{2}\right]$ between these two molecules (Fig. 4). The two molecules within the resulting six-membered ring are related to each other by a twofold rotation axis.

## 4. Synthesis and crystallization

In a typical experiment, $\alpha$-benzoinoxime was refluxed with di-$n$-butyltin oxide, $\mathrm{C}_{8} \mathrm{H}_{18} \mathrm{OSn}$, in ethanol for 2.5 h . Single crystals of the title compound suitable for X-ray diffraction were obtained from the ethanolic solution layered with $n$-hexane.

## 5. Refinement details

Crystal data, data collection and structure refinement details are summarized in Table 3. All H atoms were clearly identified in difference Fourier syntheses. Those of the carbon skeleton were calculated assuming idealized geometries and allowed to ride on the carbon atoms with $1.00 \AA$ for $s p^{3}$-hybridized and $0.95 \AA$ for aromatic H atoms, and with $U_{\text {iso }}(\mathrm{H})=1.2 U_{\text {eq }}(\mathrm{C})$. The H atoms of the two hydroxy groups were modelled with a common $\mathrm{O}-\mathrm{H}$ distance of $0.96 \AA$ before they were fixed and allowed to ride on the corresponding oxygen atom with

Table 3
Experimental details.
Crystal data Chemical formula
$M_{\mathrm{r}}$
Crystal system, space group
Temperature (K)
$a, b, c(\AA)$
$\beta\left({ }^{\circ}\right)$
$V\left(\mathrm{~A}^{3}\right)$
Z
Radiation type
$\mu\left(\mathrm{mm}^{-1}\right)$
Crystal size (mm)
Data collection
Diffractometer
Absorption correction
$T_{\text {min }}, T_{\text {max }}$
No. of measured, independent and observed $[I>2 \sigma(I)]$ reflections
$R_{\text {int }}$
$(\sin \theta / \lambda)_{\max }\left(\AA^{-1}\right)$

## Refinement

$R\left[F^{2}>2 \sigma\left(F^{2}\right)\right], w R\left(F^{2}\right), S$
No. of reflections
No. of parameters
H -atom treatment
$\Delta \rho_{\max }, \Delta \rho_{\min }\left(\mathrm{e} \AA^{-3}\right)$
$\mathrm{C}_{14} \mathrm{H}_{13} \mathrm{NO}_{2}$
227.25

Monoclinic, C2/c
100
24.1434 (9), 10.5348 (4), 8.9006 (4)
93.042 (2)
2260.64 (16)

8
Mo $K \alpha$
0.09
$0.37 \times 0.32 \times 0.11$

## Bruker APEXII CCD

Multi-scan (SADABS; Bruker, 2009)
0.968, 0.990

50071, 2005, 1765
0.043
0.595
$0.036,0.088,1.08$
2005
157
H -atom parameters constrained
$0.22,-0.18$

Computer programs: APEX2 (Bruker, 2009), SAINT (Bruker, 2009), SHELXS97 (Sheldrick, 2008), SHELXL2014 (Sheldrick, 2015), DIAMOND (Brandenburg, 2006) and Mercury (Macrae et al., 2008).
$U_{\text {iso }}(\mathrm{H})=1.2 U_{\text {eq }}(\mathrm{O})$. Disorder of the hydroxy group attached to C 2 was taken into account reducing the site occupancy of both H atoms to one-half. This suggestion was confirmed by difference-Fourier maps that clearly showed both positions.

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## supporting information

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## Crystal structure of ( $E$ )-2-hydroxy-1,2-diphenylethan-1-one oxime

Hans Reuter and Coco K. Y. A. Okio

## Computing details

Data collection: APEX2 (Bruker, 2009); cell refinement: SAINT (Bruker, 2009); data reduction: SAINT (Bruker, 2009); program(s) used to solve structure: SHELXS97 (Sheldrick, 2008); program(s) used to refine structure: SHELXL2014 (Sheldrick, 2015); molecular graphics: DIAMOND (Brandenburg, 2006) and Mercury (Macrae et al., 2006); software used to prepare material for publication: SHELXL2014 (Sheldrick, 2015).

## (E)-2-Hydroxy-1,2-diphenylethan-1-one oxime

## Crystal data

$\mathrm{C}_{14} \mathrm{H}_{13} \mathrm{NO}_{2}$
$M_{r}=227.25$
Monoclinic, $C 2 / c$
$a=24.1434$ (9) $\AA$
$b=10.5348(4) \AA$
$c=8.9006$ (4) $\AA$
$\beta=93.042(2)^{\circ}$
$V=2260.64(16) \AA^{3}$
$Z=8$

## Data collection

Bruker APEXII CCD
diffractometer
$\varphi$ and $\omega$ scans
Absorption correction: multi-scan
(SADABS; Bruker, 2009)
$T_{\text {min }}=0.968, T_{\text {max }}=0.990$
50071 measured reflections

## Refinement

Refinement on $F^{2}$
Least-squares matrix: full
$R\left[F^{2}>2 \sigma\left(F^{2}\right)\right]=0.036$
$w R\left(F^{2}\right)=0.088$
$S=1.08$
2005 reflections
157 parameters
0 restraints

$$
F(000)=960
$$

$D_{\mathrm{x}}=1.335 \mathrm{Mg} \mathrm{m}^{-3}$
Mo $K \alpha$ radiation, $\lambda=0.71073 \AA$
Cell parameters from 1932 reflections
$\theta=3.1-24.4^{\circ}$
$\mu=0.09 \mathrm{~mm}^{-1}$
$T=100 \mathrm{~K}$
Block, colourless
$0.37 \times 0.32 \times 0.11 \mathrm{~mm}$

2005 independent reflections
1765 reflections with $I>2 \sigma(I)$
$R_{\text {int }}=0.043$
$\theta_{\text {max }}=25.0^{\circ}, \theta_{\text {min }}=2.1^{\circ}$
$h=-28 \rightarrow 28$
$k=-12 \rightarrow 12$
$l=-10 \rightarrow 10$

Hydrogen site location: mixed
H -atom parameters constrained
$w=1 /\left[\sigma^{2}\left(F_{o}{ }^{2}\right)+(0.0361 P)^{2}+2.1083 P\right]$
where $P=\left(F_{0}^{2}+2 F_{\mathrm{c}}^{2}\right) / 3$
$(\Delta / \sigma)_{\text {max }}<0.001$
$\Delta \rho_{\text {max }}=0.22 \mathrm{e} \AA^{-3}$
$\Delta \rho_{\text {min }}=-0.18$ e $\AA^{-3}$

## 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.

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

|  | $x$ | $y$ | $z$ | $U_{\text {iso }} * / U_{\text {eq }}$ | Occ. $(<1)$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| O1 | 0.43413 (4) | 0.17890 (9) | 0.80605 (10) | 0.0241 (2) |  |
| H1 | 0.4693 | 0.1776 | 0.8619 | 0.029 (3)* |  |
| C1 | 0.41037 (6) | 0.14256 (12) | 0.55975 (15) | 0.0203 (3) |  |
| N1 | 0.44993 (5) | 0.16404 (11) | 0.65751 (12) | 0.0214 (3) |  |
| O2 | 0.48687 (4) | 0.10235 (9) | 0.40164 (11) | 0.0252 (3) |  |
| H3 | 0.4938 | 0.0231 | 0.4524 | 0.029 (3)* | 0.5 |
| H4 | 0.4959 | 0.0973 | 0.2981 | 0.029 (3)* | 0.5 |
| C2 | 0.42878 (6) | 0.12695 (13) | 0.39955 (15) | 0.0217 (3) |  |
| H2 | 0.4086 | 0.0531 | 0.3519 | 0.029 (3)* |  |
| C11 | 0.35015 (6) | 0.13354 (13) | 0.58375 (15) | 0.0210 (3) |  |
| C12 | 0.31705 (6) | 0.04617 (14) | 0.50248 (16) | 0.0253 (3) |  |
| H12 | 0.3332 | -0.0072 | 0.4307 | 0.0306 (19)* |  |
| C13 | 0.26100 (6) | 0.03627 (15) | 0.52511 (17) | 0.0302 (4) |  |
| H13 | 0.2391 | -0.0245 | 0.4700 | 0.0306 (19)* |  |
| C14 | 0.23671 (6) | 0.11432 (16) | 0.62737 (17) | 0.0311 (4) |  |
| H14 | 0.1982 | 0.1071 | 0.6432 | 0.0306 (19)* |  |
| C15 | 0.26863 (6) | 0.20283 (15) | 0.70637 (17) | 0.0283 (3) |  |
| H15 | 0.2518 | 0.2576 | 0.7755 | 0.0306 (19)* |  |
| C16 | 0.32497 (6) | 0.21281 (14) | 0.68598 (16) | 0.0247 (3) |  |
| H16 | 0.3466 | 0.2738 | 0.7417 | 0.0306 (19)* |  |
| C21 | 0.41529 (5) | 0.24451 (14) | 0.30732 (15) | 0.0223 (3) |  |
| C22 | 0.43786 (6) | 0.36078 (15) | 0.34820 (17) | 0.0292 (3) |  |
| H22 | 0.4627 | 0.3671 | 0.4344 | 0.040 (2)* |  |
| C23 | 0.42453 (7) | 0.46786 (16) | 0.2644 (2) | 0.0373 (4) |  |
| H23 | 0.4397 | 0.5478 | 0.2940 | 0.040 (2)* |  |
| C24 | 0.38915 (7) | 0.45886 (17) | 0.13770 (19) | 0.0402 (4) |  |
| H24 | 0.3797 | 0.5327 | 0.0807 | 0.040 (2)* |  |
| C25 | 0.36771 (7) | 0.34293 (18) | 0.09425 (18) | 0.0404 (4) |  |
| H25 | 0.3440 | 0.3362 | 0.0059 | 0.040 (2)* |  |
| C26 | 0.38058 (6) | 0.23623 (16) | 0.17912 (17) | 0.0312 (4) |  |
| H26 | 0.3654 | 0.1564 | 0.1491 | 0.040 (2)* |  |

Atomic displacement parameters $\left(\AA^{2}\right)$

|  | $U^{11}$ | $U^{22}$ | $U^{33}$ | $U^{12}$ | $U^{13}$ | $U^{23}$ |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| O1 | $0.0247(5)$ | $0.0303(6)$ | $0.0176(5)$ | $0.0021(4)$ | $0.0046(4)$ | $-0.0010(4)$ |
| C1 | $0.0231(7)$ | $0.0159(7)$ | $0.0221(7)$ | $0.0033(5)$ | $0.0038(5)$ | $0.0023(5)$ |
| N1 | $0.0240(6)$ | $0.0217(6)$ | $0.0189(6)$ | $0.0021(5)$ | $0.0057(5)$ | $-0.0002(5)$ |
| O2 | $0.0211(5)$ | $0.0259(5)$ | $0.0292(5)$ | $0.0051(4)$ | $0.0078(4)$ | $0.0033(4)$ |


| C2 | $0.0185(7)$ | $0.0234(7)$ | $0.0236(7)$ | $0.0017(5)$ | $0.0041(5)$ | $-0.0008(6)$ |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| C11 | $0.0230(7)$ | $0.0209(7)$ | $0.0194(7)$ | $0.0025(5)$ | $0.0037(5)$ | $0.0051(5)$ |
| C12 | $0.0273(8)$ | $0.0258(8)$ | $0.0230(7)$ | $0.0027(6)$ | $0.0029(6)$ | $0.0014(6)$ |
| C13 | $0.0256(8)$ | $0.0347(9)$ | $0.0301(8)$ | $-0.0045(6)$ | $-0.0001(6)$ | $0.0031(7)$ |
| C14 | $0.0224(8)$ | $0.0421(9)$ | $0.0292(8)$ | $0.0007(7)$ | $0.0052(6)$ | $0.0090(7)$ |
| C15 | $0.0266(8)$ | $0.0332(8)$ | $0.0259(8)$ | $0.0060(6)$ | $0.0084(6)$ | $0.0035(6)$ |
| C16 | $0.0268(8)$ | $0.0245(7)$ | $0.0232(7)$ | $0.0024(6)$ | $0.0043(6)$ | $0.0022(6)$ |
| C21 | $0.0209(7)$ | $0.0265(8)$ | $0.0203(7)$ | $0.0057(6)$ | $0.0081(5)$ | $0.0002(6)$ |
| C22 | $0.0269(8)$ | $0.0314(8)$ | $0.0297(8)$ | $0.0012(6)$ | $0.0051(6)$ | $0.0027(7)$ |
| C23 | $0.0397(9)$ | $0.0274(9)$ | $0.0465(10)$ | $0.0026(7)$ | $0.0177(8)$ | $0.0054(7)$ |
| C24 | $0.0503(10)$ | $0.0395(10)$ | $0.0325(9)$ | $0.0225(8)$ | $0.0189(8)$ | $0.0163(8)$ |
| C25 | $0.0478(10)$ | $0.0527(11)$ | $0.0206(8)$ | $0.0244(9)$ | $0.0007(7)$ | $0.0016(7)$ |
| C26 | $0.0346(8)$ | $0.0353(9)$ | $0.0239(8)$ | $0.0095(7)$ | $0.0028(6)$ | $-0.0052(7)$ |

Geometric parameters ( $A,{ }^{\circ}$ )

| $\mathrm{O} 1-\mathrm{N} 1$ | 1.404 (1) | C14-C15 | 1.378 (2) |
| :---: | :---: | :---: | :---: |
| $\mathrm{O} 1-\mathrm{H} 1$ | 0.9600 | C14-H14 | 0.9500 |
| $\mathrm{C} 1-\mathrm{N} 1$ | 1.278 (2) | C15-C16 | 1.386 (2) |
| C1-C11 | 1.484 (2) | C15-H15 | 0.9500 |
| $\mathrm{C} 1-\mathrm{C} 2$ | 1.525 (2) | C16-H16 | 0.9500 |
| $\mathrm{O} 2-\mathrm{C} 2$ | 1.425 (2) | C21-C22 | 1.381 (2) |
| O 2 - H 3 | 0.9600 | C21-C26 | 1.382 (2) |
| $\mathrm{O} 2-\mathrm{H} 4$ | 0.9600 | C22-C23 | 1.381 (2) |
| C2-C21 | 1.512 (2) | C22-H22 | 0.9500 |
| C2-H2 | 1.0000 | C23-C24 | 1.382 (3) |
| C11-C12 | 1.396 (2) | C23-H23 | 0.9500 |
| C11-C16 | 1.398 (2) | C24-C25 | 1.374 (3) |
| C12-C13 | 1.383 (2) | C24-H24 | 0.9500 |
| C12-H12 | 0.9500 | C25-C26 | 1.380 (2) |
| C13-C14 | 1.380 (2) | C25-H25 | 0.9500 |
| C13-H13 | 0.9500 | C26-H26 | 0.9500 |
| N1-O1-H1 | 102.1 | C13-C14-H14 | 120.2 |
| N1-C1-C11 | 128.03 (12) | C14-C15-C16 | 120.70 (14) |
| N1-C1-C2 | 114.3 (1) | C14-C15-H15 | 119.6 |
| $\mathrm{C} 11-\mathrm{C} 1-\mathrm{C} 2$ | 117.7 (1) | C16-C15-H15 | 119.6 |
| $\mathrm{C} 1-\mathrm{N} 1-\mathrm{O} 1$ | 115.5 (1) | C15-C16-C11 | 120.25 (14) |
| $\mathrm{C} 2-\mathrm{O} 2-\mathrm{H} 3$ | 108.1 | C15-C16-H16 | 119.9 |
| $\mathrm{C} 2-\mathrm{O} 2-\mathrm{H} 4$ | 105.7 | C11-C16-H16 | 119.9 |
| H3-O2-H4 | 111.2 | C22-C21-C26 | 119.12 (14) |
| $\mathrm{O} 2-\mathrm{C} 2-\mathrm{C} 21$ | 109.84 (11) | C22-C21-C2 | 120.85 (13) |
| $\mathrm{O} 2-\mathrm{C} 2-\mathrm{C} 1$ | 110.1 (1) | C26-C21-C2 | 120.02 (13) |
| $\mathrm{C} 21-\mathrm{C} 2-\mathrm{C} 1$ | 110.75 (11) | C23-C22-C21 | 120.32 (15) |
| $\mathrm{O} 2-\mathrm{C} 2-\mathrm{H} 2$ | 108.7 | $\mathrm{C} 23-\mathrm{C} 22-\mathrm{H} 22$ | 119.8 |
| C21-C2-H2 | 108.7 | $\mathrm{C} 21-\mathrm{C} 22-\mathrm{H} 22$ | 119.8 |
| $\mathrm{C} 1-\mathrm{C} 2-\mathrm{H} 2$ | 108.7 | C22-C23-C24 | 120.07 (16) |
| C12-C11-C16 | 118.33 (13) | C22-C23-H23 | 120.0 |


| C12-C11-C1 | $120.41(12)$ | C24-C23-H23 | 120.0 |
| :--- | :--- | :--- | :--- |
| C16-C11-C1 | $121.26(13)$ | C25-C24-C23 | $119.88(15)$ |
| C13-C12-C11 | $120.81(13)$ | C25-C24-H24 | 120.1 |
| C13-C12-H12 | 119.6 | C23-C24-H24 | 120.1 |
| C11-C12-H12 | 119.6 | C24-C25-C26 | $119.95(16)$ |
| C14-C13-C12 | $120.31(14)$ | C24-C25-H25 | 120.0 |
| C14-C13-H13 | 119.8 | C26-C25-H25 | 120.0 |
| C12-C13-H13 | 119.8 | C25-C26-C21 | $120.62(16)$ |
| C15-C14-C13 | $119.58(14)$ | C25-C26-H26 | 119.7 |
| C15-C14-H14 | 120.2 | C21-C26-H26 | 119.7 |
| C11-C1-N1-O1 | $1.1(2)$ |  |  |
| C2-C1-N1-O1 | $179.96(10)$ | C14-C15-C16-C11 | $0.5(2)$ |
| N1-C1-C2-O2 | $17.34(16)$ | C1-C11-C16-C15 | $0.8(2)$ |
| C11-C1-C2-O2 | $-163.64(11)$ | O2-C2-C21-C22 | $179.89(13)$ |
| N1-C1-C2-C21 | $-104.31(14)$ | C1-C2-C21-C22 | $-61.09(16)$ |
| C11-C1-C2-C21 | $74.72(15)$ | O2-C2-C21-C26 | $60.68(16)$ |
| N1-C1-C11-C12 | $-143.31(15)$ | C1-C2-C21-C26 | $-1120.23(14)$ |
| C2-C1-C11-C12 | $37.82(18)$ | C26-C21-C22-C23 | $1.9(2)$ |
| N1-C1-C11-C16 | $37.6(2)$ | C2-C21-C22-C23 | $-178.96(13)$ |
| C2-C1-C11-C16 | $-141.28(13)$ | C21-C22-C23-C24 | $-1.0(2)$ |
| C16-C11-C12-C13 | $-1.5(2)$ | C22-C23-C24-C25 | $-0.7(2)$ |
| C1-C11-C12-C13 | $179.39(13)$ | C23-C24-C25-C26 | $1.4(2)$ |
| C11-C12-C13-C14 | $0.9(2)$ | C24-C25-C26-C21 | $-0.5(2)$ |
| C12-C13-C14-C15 | $0.3(2)$ | C22-C21-C26-C25 | $-1.2(2)$ |
| C13-C14-C15-C16 | $-1.1(2)$ | C2-C21-C26-C25 | $179.69(13)$ |

Hydrogen-bond geometry ( $A,{ }^{\circ}$ )

| $D-\mathrm{H} \cdots A$ | $D-\mathrm{H}$ | $\mathrm{H} \cdots A$ | $D \cdots A$ | $D-\mathrm{H} \cdots A$ |
| :--- | :--- | :--- | :--- | :--- |
| $\mathrm{O} 1 — \mathrm{H} 1 \cdots \mathrm{~N}^{\mathrm{i}}$ | 0.96 | 1.97 | $2.805(2)$ | 144 |
| $\mathrm{O} 2-\mathrm{H} 3 \cdots \mathrm{O}^{2 i}$ | 0.96 | 1.89 | $2.829(2)$ | 164 |
| $\mathrm{O} 2-\mathrm{H} 4 \cdots \mathrm{O}^{2 i i}$ | 0.96 | 1.85 | $2.806(2)$ | 175 |

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

