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Crystal structure of 1,4-diethoxy-9,10anthraquinone

Chitoshi Kitamura,^a* Sining Li,^a Munenori Takehara,^a Yoshinori Inoue,^a Katsuhiko Ono^b and Takeshi Kawase^c

^aDepartment of Materials Science, School of Engineering, The University of Shiga Prefecture, 2500 Hassaka-cho, Hikone, Shiga 522-8533, Japan, ^bDepartment of Materials Science and Engineering, Graduate School of Engineering, Nagoya Institute of Technology, Gokiso, Showa-ku, Nagoya, Aichi 466-8555, Japan, and ^cDepartment of Applied Chemistry, Graduate School of Engineering, University of Hyogo, 2167 Shosha, Himeji, Hyogo 671-2280, Japan. *Correspondence e-mail: kitamura.c@mat.usp.ac.jp

Received 16 June 2015; accepted 22 June 2015

Edited by H. Ishida, Okayama University, Japan

The asymmetric unit of the title compound, $C_{18}H_{16}O_4$, contains two crystallographically independent molecules. The anthraquinone ring systems are slightly bent with dihedral angles of 2.33 (8) and $13.31 (9)^{\circ}$ between the two terminal benzene rings. In the crystal, the two independent molecules adopt slipped-parallel π -overlap with an average interplanar distance of 3.45 Å, forming a dimer; the centroid-centroid distances of the π - π interactions are 3.6659 (15)-3.8987 (15) Å. The molecules are also linked by $C-H \cdots O$ interactions, forming a tape structure along the a-axis direction. The crystal packing is characterized by a dimerherringbone pattern.

Keywords: crystal structure; 9,10-anthraquinone; crystallographically independent molecules; $\pi - \pi$ interactions; C—H···O interactions.

CCDC reference: 1008606

1. Related literature

For synthesis of alkoxy-substituted 9,10-anthraquinones, see: Kitamura et al. (2004). For background information on substitution effects of alkoxy-substituted 9,10-anthraquinones, see; Ohta et al. (2012). For related structures of 1,4-dipropoxy-9,10-anthraquinone polymorphs, see: Kitamura et al. (2015).



V = 2910.4 (4) Å³

Mo $K\alpha$ radiation

 $0.56 \times 0.40 \times 0.36$ mm

6645 independent reflections

3129 reflections with $I > 2\sigma(I)$

 $\mu = 0.10 \text{ mm}^-$

T = 223 K

 $R_{\rm int} = 0.045$

Z = 8

2. Experimental

2.1. Crystal data

 $C_{18}H_{16}O_4$ $M_r = 296.31$ Monoclinic, $P2_1/c$ a = 13.5514 (11) Åb = 14.7204 (11) Åc = 14.5905 (10) Å $\beta = 90.604 \ (3)^{\circ}$

2.2. Data collection

Rigaku R-AXIS RAPID diffractometer 27699 measured reflections

2.3. Refinement

$R[F^2 > 2\sigma(F^2)] = 0.076$	397 parameters
$wR(F^2) = 0.273$	H-atom parameters constrained
S = 0.93	$\Delta \rho_{\rm max} = 0.27 \ {\rm e} \ {\rm \AA}^{-3}$
6645 reflections	$\Delta \rho_{\rm min} = -0.48 \ {\rm e} \ {\rm \AA}^{-3}$

Table 1

Hydrogen-bond geometry (Å, °).

$D - H \cdots A$	$D-\mathrm{H}$	$H \cdot \cdot \cdot A$	$D \cdots A$	$D - \mathbf{H} \cdots \mathbf{A}$
$C8A - H8A \cdots O3B$	0.94	2.48	3.234 (3)	137
$C8B - H8B \cdots O3A$	0.94	2.55	3.304 (4)	137
$C11A - H11A \cdots O4B^{i}$	0.94	2.60	3.325 (3)	135
$C11B - H11B \cdots O4A^{ii}$	0.94	2.46	3.199 (4)	135

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

Data collection: PROCESS-AUTO (Rigaku, 1998); cell refinement: PROCESS-AUTO; data reduction: PROCESS-AUTO; program(s) used to solve structure: SIR2004 (Burla et al., 2005); program(s) used to refine structure: SHELXL2014 (Sheldrick, 2015); molecular graphics: ORTEP-3 for Windows (Farrugia, 2012) and Mercury (Macrae et al., 2008); software used to prepare material for publication: WinGX (Farrugia, 2012).

Acknowledgements

This work was supported by a Grant-in-Aid for Scientific Research (C) (No. 15 K05482) from the JSPS.

Supporting information for this paper is available from the IUCr electronic archives (Reference: IS5404).

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supporting information

Acta Cryst. (2015). E71, o504–o505 [doi:10.1107/S2056989015011901]

Crystal structure of 1,4-diethoxy-9,10-anthraquinone

Chitoshi Kitamura, Sining Li, Munenori Takehara, Yoshinori Inoue, Katsuhiko Ono and Takeshi Kawase

S1. Comment

9,10-Anthraquinone is an important framework as a dye. Various kinds of hydroxy-substituted anthraquinone dyes have been manufactured. However, there were little reports on alkoxy-substituted anthraquinone. In recent years, we presented the effects of the alkoxy substitution on the optical properties of 2,6-dialkoxy and 2,3,6,7-tetraalkoxy derivatives in solution as well as in the solid state (Ohta *et al.*, 2012). Very recently, we have reported crystal structures of two polymorphs of 1,4-dipropoxy-9,10-anthraquinone, which contained red and yellow solids (Kitamura *et al.*, 2015). The red crystal exhibited an anti-parallel arrangement along the stacking direction. On the other hand, the yellow crystal showed a slipped-parallel arrangement. To search the effect of alkyl chain length on molecular packing, we prepared the title compound, 1,4-diethoxy-9,10-anthraquinone, (I). In this paper, we present the crystal structure of (I).

The molecular structure of (I) is shown in Fig. 1. Two crystallographically independent molecules were found in the asymmetric unit, although the two molecules had almost the same molecular structure. There was a difference in planarity between the two molecules. Thus, the anthraquinone framework was slightly bent at the central quinone ring. For example, the dihedral angle between the two terminal benzene rings in the anthraquinone was 2.33 (8)° for one molecule and 13.31 (9)° for the other. The packing structure displays a dimer-herringbone pattern (Fig. 2), which is completely different from those of 1,4-dipropoxy-9,10-anthraquinone polymorphs (Kitamura *et al.*, 2015). In the dimer part, the two molecules adopt slipped-parallel π -stack with an average interplanar distance of 3.45 Å, which would result in a yellow color in the solid state. The crystal structure is also stabilized by C—H…O interactions along the lateral direction of molecules (Fig. 3).

S2. Experimental

The title compound was prepared according to our previously reported method (Kitamura *et al.*, 2004). A mixture of 1,4hydrooxy-9,10-anthraquinone (2.20 g, 9.16 mmol), K₂CO₃ (2.51 g, 18.1 mmol), ethyl *p*-toluenesulfonate (5.02 g, 25.1 mmol) in *o*-dichlorobenzene (15 ml) was heated at reflux for 3 h under N₂ gas. After cooling to room temperature, water (65 ml) was added to the reaction mixture. Then, the resulting solid was filtered off and washed with hexane to give the title compound (2.37 g, 87% yield) as a yellow solid. Single crystals suitable for X-ray analysis were obtained by slow evaporation from a CH₂Cl₂ solution (*m.p.* 172–175 °C). Elemental analysis for C₁₈H₁₆O₄: C 72.96, H 5.44. Found: C 72.75, H 5.51. TOF-MS(EI): *m/z* Calcd C₁₈H₁₆O₄: 296.1049. Found: 296.1074.

S3. Refinement

All the H atoms were positioned geometrically and refined using a riding model with C—H bonds of 0.94 Å, 0.98 Å, and 0.97 Å for aromatic, methylene and methyl groups, respectively, and $U_{iso}(H) = 1.2U_{eq}(C)$ [$U_{iso}(H) = 1.5U_{eq}(C)$ for methyl H atoms].



Figure 1

The asymmetric unit of the title compound, showing the atomic numbering and 40% probability displacement ellipsoids.



Figure 2

A packing diagram of the title compound viewed down the *a* axis, showing a dimer-herringbone pattern. Hydrogen atoms are omitted for clarity.



Figure 3

A packing diagram of the title compound, showing C—H…O interactions (blue lines).

1,4-Diethoxy-9,10-anthraquinone

Crystal data	
C ₁₈ H ₁₆ O ₄ $M_r = 296.31$ Monoclinic, $P2_1/c$ Hall symbol: -P 2ybc a = 13.5514 (11) Å b = 14.7204 (11) Å c = 14.5905 (10) Å $\beta = 90.604 (3)^{\circ}$ $V = 2910.4 (4) \text{ Å}^3$ Z = 8	F(000) = 1248 $D_x = 1.352 \text{ Mg m}^{-3}$ Mo K α radiation, $\lambda = 0.71073 \text{ Å}$ Cell parameters from 11158 reflections $\theta = 3-27.5^{\circ}$ $\mu = 0.10 \text{ mm}^{-1}$ T = 223 K Prism, orange $0.56 \times 0.40 \times 0.36 \text{ mm}$
Data collection Rigaku R-AXIS RAPID diffractometer Radiation source: fine-focus sealed x-ray tube Graphite monochromator Detector resolution: 10 pixels mm ⁻¹ ω scans 27699 measured reflections	6645 independent reflections 3129 reflections with $I > 2\sigma(I)$ $R_{int} = 0.045$ $\theta_{max} = 27.5^{\circ}, \theta_{min} = 3.0^{\circ}$ $h = -17 \rightarrow 17$ $k = -19 \rightarrow 19$ $l = -16 \rightarrow 18$

Refinement

Refinement on F^2	0 constraints
Least-squares matrix: full	H-atom parameters constrained
$R[F^2 > 2\sigma(F^2)] = 0.076$	$w = 1/[\sigma^2(F_o^2) + (0.1824P)^2]$
$wR(F^2) = 0.273$	where $P = (F_o^2 + 2F_c^2)/3$
S = 0.93	$(\Delta/\sigma)_{\rm max} < 0.001$
6645 reflections	$\Delta ho_{ m max} = 0.27 \ m e \ m \AA^{-3}$
397 parameters	$\Delta \rho_{\rm min} = -0.48$ e Å ⁻³
0 restraints	
0 restraints	

Special details

Experimental. ¹H-NMR: δ 1.56 (t, *J* = 7.0 Hz, 6H), 4.20 (q, *J* = 7.0 Hz, 4H), 7.32 (s, 2H), 7.69–7.72 (m, 2H), 8.17–8.19 (m, 2H); ¹³C-NMR: δ 14.9, 66.0, 122.1, 123.4, 126.4, 133.2, 134.2, 153.6, 183.3.

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 $(Å^2)$

	x	У	Ζ	$U_{\rm iso}$ */ $U_{\rm eq}$
C1A	0.3813 (2)	0.68438 (17)	0.35528 (16)	0.0548 (6)
C2A	0.3271 (2)	0.71337 (18)	0.27977 (17)	0.0611 (7)
H2A	0.3605	0.7327	0.2271	0.073*
C3A	0.2261 (2)	0.71464 (18)	0.27990 (16)	0.0596 (7)
H3A	0.1918	0.7347	0.2274	0.072*
C4A	0.1731 (2)	0.68680 (17)	0.35617 (16)	0.0546 (6)
C5A	0.22559 (19)	0.65664 (16)	0.43427 (16)	0.0518 (6)
C6A	0.1713 (2)	0.63214 (19)	0.51927 (18)	0.0611 (7)
C7A	0.22803 (19)	0.59444 (17)	0.59796 (15)	0.0527 (6)
C8A	0.1770 (2)	0.56507 (18)	0.67540 (17)	0.0632 (7)
H8A	0.1078	0.5678	0.6766	0.076*
C9A	0.2293 (2)	0.5320 (2)	0.75003 (17)	0.0707 (8)
H9A	0.1952	0.5121	0.8021	0.085*
C10A	0.3301 (2)	0.5279 (2)	0.74903 (18)	0.0726 (8)
H10A	0.3647	0.5057	0.8005	0.087*
C11A	0.3814 (2)	0.5564 (2)	0.67253 (18)	0.0681 (8)
H11A	0.4507	0.553	0.6717	0.082*
C12A	0.3296 (2)	0.59013 (18)	0.59665 (17)	0.0572 (6)
C13A	0.3855 (2)	0.6237 (2)	0.5163 (2)	0.0734 (9)
C14A	0.32994 (19)	0.65421 (16)	0.43355 (16)	0.0530 (6)
C15A	0.5325 (2)	0.7171 (2)	0.2792 (2)	0.0782 (9)
H15A	0.5119	0.7793	0.265	0.094*
H15B	0.5174	0.6787	0.226	0.094*
C16A	0.6399 (2)	0.7146 (2)	0.2998 (2)	0.0814 (9)
H16A	0.676	0.7365	0.2471	0.122*
H16B	0.6542	0.7531	0.3523	0.122*
H16C	0.6598	0.6527	0.3134	0.122*
C17A	0.0200 (2)	0.7234 (2)	0.28108 (18)	0.0672 (7)

H17A	0.0363	0.6897	0.2253	0.081*
H17B	0.0373	0.7874	0.2719	0.081*
C18A	-0.0867 (2)	0.7146 (2)	0.3010(2)	0.0787 (9)
H18A	-0.1251	0.7386	0.25	0.118*
H18B	-0.1029	0.651	0.3099	0.118*
H18C	-0.1019	0.7483	0.3562	0.118*
O1A	0.48108 (14)	0.68464 (14)	0.35759 (12)	0.0673 (5)
O2A	0.07320 (14)	0.68735 (14)	0.35765 (12)	0.0670 (5)
O3A	0.08375 (17)	0.6462 (2)	0.52801 (15)	0.1020 (9)
O4A	0.47401 (18)	0.6270 (3)	0.5216 (2)	0.1549 (16)
C1B	-0.31908(18)	0.42295(17)	0.93060(15)	0.0515 (6)
C2B	-0.2634(2)	0.3807(2)	0.99876 (16)	0.0600(7)
H2B	-0.296	0.3526	1 0478	0.072*
C3B	-0.1627(2)	0.37894(19)	0.99648 (16)	0.0582(7)
H3B	-0.1276	0.3486	1 0432	0.07*
C4B	-0.11091(18)	0.42118 (16)	0.92621(15)	0.0507 (6)
C5B	-0.16423(18)	0.46877(16)	0.85830(14)	0.0207(0) 0.0477(6)
C6B	-0.11215(18)	0.10077(10) 0.51742(17)	0.03030(11) 0.78342(15)	0.0517(6)
C7B	-0.17213(18)	0.54479(16)	0.70312(15) 0.70136(15)	0.0317(0) 0.0495(6)
C8B	-0.1216(2)	0.57178(19)	0.62221(17)	0.0195(0)
H8B	-0.0523	0.5707	0.6203	0.0050(7)
C9B	-0.1754(2)	0.5707	0.54668(17)	0.0683 (8)
H9B	-0.1426	0.6169	0.4928	0.082*
C10B	-0.2760(2)	0.6109	0.54962 (17)	0.062
H10B	-0.312	0.6241	0.498	0.082*
C11B	-0.3252(2)	0.57867 (18)	0.62810(17)	0.002 0.0634(7)
H11B	-0.3944	0.5823	0.63	0.0051(7)
C12B	-0.27238(18)	0.54778 (16)	0.05	0.0499 (6)
C13B	-0.32523(18)	0.52046 (17)	0.78847(15)	0.0520(6)
C14B	-0.26902(17)	0.32010(17) 0.47017(16)	0.86019 (14)	0.0520(0) 0.0474(5)
C15B	-0.4706(2)	0.17017(10) 0.3711(2)	0.00019(11) 0.99707(18)	0.0672(8)
H15C	-0.462	0.4013	1.0565	0.081*
H15D	-0.4444	0.3092	1.0022	0.081*
C16B	-0.5768(2)	0.3682(2)	0.97143(19)	0.001 0.0714(8)
H16D	-0.6129	0.3351	1 0178	0.107*
H16E	-0.6022	0.4296	0.9669	0.107*
H16F	-0.5847	0.3378	0.9128	0.107*
C17B	0.04137(19)	0.36059 (19)	0.98395(17)	0.0605 (7)
H17C	0.0159	0.2983	0.9816	0.0005 (7)
H17D	0.0335	0.3839	1 0464	0.073*
C18B	0.0355 0.1478(2)	0.3621(2)	0.95790(19)	0.075 0.0681 (7)
H18D	0.1855	0.3244	1 0001	0.102*
H18E	0.1546	0.3388	0.8961	0.102*
H18F	0.1722	0.424	0.9606	0.102*
O1B	-0.41883(13)	0.42021(13)	0.92745(11)	0.0609 (5)
02B	-0.01133(12)	0.41697(12)	0.92743(11) 0.91984(11)	0.0009(5)
03B	-0.02445(14)	0 53591 (16)	0.78861(12)	0.0763 (6)
04B	-0.41212(14)	0.53571(10) 0.54036(16)	0 79662 (13)	0.0702 (6)
	0.71212(17)	0.27020(10)	0.19002 (19)	0.0772(0)

Atomic displacement parameters $(Å^2)$

	U^{11}	U ²²	U ³³	U^{12}	U ¹³	U ²³
C1A	0.0605 (17)	0.0546 (14)	0.0495 (13)	-0.0036 (11)	0.0078 (11)	-0.0019 (11)
C2A	0.0725 (19)	0.0649 (16)	0.0460 (13)	-0.0059 (13)	0.0096 (12)	0.0024 (11)
C3A	0.0730 (19)	0.0623 (15)	0.0436 (13)	-0.0014 (13)	-0.0008 (12)	0.0022 (11)
C4A	0.0588 (17)	0.0563 (14)	0.0488 (13)	-0.0021 (11)	-0.0015 (11)	0.0009 (11)
C5A	0.0555 (15)	0.0521 (13)	0.0478 (13)	0.0009 (10)	0.0027 (11)	0.0039 (10)
C6A	0.0500 (16)	0.0738 (18)	0.0595 (15)	0.0006 (12)	0.0044 (12)	0.0125 (13)
C7A	0.0571 (16)	0.0554 (14)	0.0456 (13)	-0.0020 (11)	0.0035 (11)	0.0040 (10)
C8A	0.0642 (18)	0.0737 (17)	0.0519 (14)	-0.0018 (13)	0.0090 (12)	0.0057 (12)
C9A	0.085 (2)	0.0814 (19)	0.0452 (14)	-0.0047 (16)	0.0060 (13)	0.0117 (13)
C10A	0.080 (2)	0.087 (2)	0.0507 (15)	-0.0024 (16)	-0.0112 (13)	0.0135 (14)
C11A	0.0623 (18)	0.0812 (19)	0.0605 (16)	-0.0011 (13)	-0.0049 (13)	0.0164 (14)
C12A	0.0600 (17)	0.0620 (15)	0.0497 (13)	-0.0015 (12)	-0.0007 (11)	0.0084 (11)
C13A	0.0499 (17)	0.102 (2)	0.0680 (17)	-0.0035 (15)	0.0006 (13)	0.0333 (16)
C14A	0.0550 (15)	0.0549 (14)	0.0490 (13)	-0.0013 (11)	0.0026 (11)	0.0048 (10)
C15A	0.072 (2)	0.100 (2)	0.0625 (17)	-0.0060 (17)	0.0221 (15)	0.0049 (16)
C16A	0.068 (2)	0.099 (2)	0.078 (2)	-0.0059 (17)	0.0214 (16)	0.0005 (17)
C17A	0.0692 (19)	0.0800 (19)	0.0522 (14)	0.0096 (14)	-0.0100 (12)	0.0001 (13)
C18A	0.068 (2)	0.102 (2)	0.0660 (17)	0.0023 (16)	-0.0154 (14)	-0.0043 (16)
O1A	0.0570 (12)	0.0860 (13)	0.0593 (11)	-0.0029 (9)	0.0144 (9)	0.0075 (9)
O2A	0.0574 (12)	0.0872 (13)	0.0563 (10)	-0.0004 (9)	-0.0067 (8)	0.0116 (9)
O3A	0.0555 (14)	0.165 (3)	0.0855 (15)	0.0170 (14)	0.0145 (11)	0.0563 (15)
O4A	0.0496 (16)	0.289 (4)	0.126 (2)	-0.0097 (19)	-0.0050 (14)	0.127 (3)
C1B	0.0491 (14)	0.0616 (14)	0.0440 (12)	-0.0026 (11)	0.0030 (10)	0.0047 (10)
C2B	0.0609 (17)	0.0720 (17)	0.0473 (13)	-0.0048 (13)	0.0066 (12)	0.0168 (12)
C3B	0.0557 (16)	0.0688 (16)	0.0500 (13)	0.0000 (12)	-0.0007 (11)	0.0154 (12)
C4B	0.0514 (15)	0.0590 (14)	0.0416 (12)	0.0001 (11)	-0.0002 (10)	0.0020 (10)
C5B	0.0534 (14)	0.0556 (13)	0.0342 (11)	-0.0005 (10)	0.0029 (9)	0.0021 (9)
C6B	0.0455 (14)	0.0653 (15)	0.0442 (12)	-0.0025 (11)	0.0026 (10)	0.0057 (11)
C7B	0.0543 (15)	0.0537 (13)	0.0407 (11)	-0.0022 (10)	0.0020 (10)	0.0059 (10)
C8B	0.0624 (17)	0.0802 (18)	0.0483 (13)	-0.0060 (14)	0.0066 (12)	0.0134 (12)
C9B	0.075 (2)	0.085 (2)	0.0451 (13)	-0.0091 (15)	0.0059 (13)	0.0173 (13)
C10B	0.074 (2)	0.0826 (19)	0.0474 (14)	-0.0010 (15)	-0.0059 (13)	0.0200 (13)
C11B	0.0597 (17)	0.0770 (18)	0.0534 (14)	0.0034 (13)	-0.0030 (12)	0.0164 (13)
C12B	0.0546 (15)	0.0539 (13)	0.0413 (12)	0.0013 (10)	0.0028 (10)	0.0056 (10)
C13B	0.0475 (14)	0.0630 (15)	0.0455 (12)	0.0023 (11)	0.0027 (10)	0.0063 (11)
C14B	0.0511 (14)	0.0551 (13)	0.0359 (11)	0.0002 (10)	0.0036 (9)	0.0007 (9)
C15B	0.0576 (18)	0.0847 (19)	0.0596 (16)	-0.0086 (14)	0.0113 (13)	0.0203 (14)
C16B	0.0570 (18)	0.095 (2)	0.0620 (16)	-0.0103 (15)	0.0097 (13)	0.0086 (15)
C17B	0.0567 (17)	0.0672 (16)	0.0572 (14)	-0.0002 (12)	-0.0105 (12)	0.0086 (12)
C18B	0.0550 (17)	0.0804 (19)	0.0688 (17)	0.0069 (14)	-0.0070 (13)	0.0045 (14)
O1B	0.0471 (11)	0.0822 (13)	0.0535 (10)	-0.0018 (8)	0.0076 (8)	0.0159 (8)
O2B	0.0478 (11)	0.0773 (12)	0.0508 (9)	0.0029 (8)	-0.0036 (7)	0.0139 (8)
O3B	0.0512 (12)	0.1171 (17)	0.0605 (11)	-0.0142 (11)	-0.0020 (8)	0.0288 (11)
O4B	0.0525 (12)	0.1193 (17)	0.0659 (12)	0.0183 (11)	0.0086 (9)	0.0321 (11)

Geometric parameters (Å, °)

C1A—O1A	1.352 (3)	C1B—01B	1.353 (3)
C1A—C2A	1.385 (4)	C1B—C2B	1.388 (3)
C1A—C14A	1.415 (3)	C1B—C14B	1.419 (3)
C2A—C3A	1.368 (4)	C2B—C3B	1.366 (3)
C2A—H2A	0.94	C2B—H2B	0.94
C3A—C4A	1.392 (3)	C3B—C4B	1.395 (3)
СЗА—НЗА	0.94	C3B—H3B	0.94
C4A—O2A	1.354 (3)	C4B—O2B	1.355 (3)
C4A—C5A	1.409 (3)	C4B—C5B	1.407 (3)
C5A—C14A	1.415 (3)	C5B—C14B	1.421 (3)
C5A—C6A	1.493 (3)	C5B—C6B	1.490 (3)
C6A—O3A	1.212 (3)	C6B—O3B	1.221 (3)
C6A—C7A	1.483 (3)	C6B—C7B	1.481 (3)
C7A—C12A	1.378 (4)	C7B—C12B	1.387 (3)
C7A—C8A	1.399 (3)	C7B—C8B	1.393 (3)
C8A—C9A	1.381 (4)	C8B—C9B	1.379 (4)
C8A—H8A	0.94	C8B—H8B	0.94
C9A-C10A	1.367 (4)	C9B—C10B	1.366 (4)
С9А—Н9А	0.94	C9B—H9B	0.94
C10A—C11A	1.386 (4)	C10B—C11B	1.383 (4)
C10A—H10A	0.94	C10B—H10B	0.94
C11A—C12A	1.396 (4)	C11B—C12B	1.392 (3)
C11A—H11A	0.94	C11B—H11B	0.94
C12A—C13A	1.488 (4)	C12B—C13B	1.484 (3)
C13A—O4A	1.202 (3)	C13B—O4B	1.220 (3)
C13A—C14A	1.485 (4)	C13B—C14B	1.485 (3)
C15A—O1A	1.428 (3)	C15B—O1B	1.436 (3)
C15A—C16A	1.484 (4)	C15B—C16B	1.484 (4)
C15A—H15A	0.98	C15B—H15C	0.98
C15A—H15B	0.98	C15B—H15D	0.98
C16A—H16A	0.97	C16B—H16D	0.97
C16A—H16B	0.97	C16B—H16E	0.97
C16A—H16C	0.97	C16B—H16F	0.97
C17A—O2A	1.425 (3)	C17B—O2B	1.435 (3)
C17A—C18A	1.485 (4)	C17B—C18B	1.495 (4)
C17A—H17A	0.98	C17B—H17C	0.98
C17A—H17B	0.98	C17B—H17D	0.98
C18A—H18A	0.97	C18B—H18D	0.97
C18A—H18B	0.97	C18B—H18E	0.97
C18A—H18C	0.97	C18B—H18F	0.97
O1A—C1A—C2A	122.8 (2)	O1B—C1B—C2B	123.1 (2)
01A—C1A—C14A	118.8 (2)	O1B—C1B—C14B	118.4 (2)
C2A—C1A—C14A	118.5 (3)	C2B—C1B—C14B	118.5 (2)
C3A—C2A—C1A	121.7 (2)	C3B—C2B—C1B	121.8 (2)
C3A—C2A—H2A	119.2	C3B—C2B—H2B	119.1

C1A—C2A—H2A	119.2	C1B—C2B—H2B	1191
$C^2A - C^3A - C^4A$	121 4 (2)	C^{2B} C^{3B} C^{4B}	1214(2)
$C_2 \Delta - C_3 \Delta - H_3 \Delta$	110.3	C^{2B} C^{3B} H^{3B}	110.3
C_{4A} C_{3A} H_{3A}	110.3	CAB C3B H3B	110.3
$O_{A} C_{A} C_{A} C_{A}$	122 3 (2)	$O^{2}B C^{4}B C^{3}B$	117.5 122.6(2)
$O_{2A} = C_{4A} = C_{5A}$	122.3(2) 1100(2)	$O_{2B} = C_{4B} = C_{5B}$	122.0(2)
$C_{2A} = C_{4A} = C_{5A}$	119.0(2) 118.6(2)	C_{2D} C_{4D} C_{5D}	118.0(2)
$C_{AA} = C_{AA} = C_{AA} = C_{AA}$	110.0(3)	C_{3B} C_{4B} C_{5B} C_{14B}	110.7(2)
C4A = C5A = C14A	119.9 (2)	C4D = C5D = C14D	120.0(2)
$C_{4A} = C_{5A} = C_{6A}$	119.9 (2)	$C_{4B} = C_{3B} = C_{0B}$	120.8(2)
C14A - C5A - C6A	120.1 (2)		119.24 (19)
03A—C6A—C/A	118.9 (2)	03B—C6B—C7B	119.8 (2)
03A—C6A—C5A	122.5 (2)	O3B—C6B—C5B	122.0 (2)
C7A—C6A—C5A	118.5 (2)	C7B—C6B—C5B	118.2 (2)
C12A—C7A—C8A	119.9 (2)	C12B—C7B—C8B	119.8 (2)
C12A—C7A—C6A	121.1 (2)	C12B—C7B—C6B	120.4 (2)
C8A—C7A—C6A	119.0 (2)	C8B—C7B—C6B	119.8 (2)
C9A—C8A—C7A	119.5 (3)	C9B—C8B—C7B	119.9 (3)
C9A—C8A—H8A	120.3	C9B—C8B—H8B	120
C7A—C8A—H8A	120.3	C7B—C8B—H8B	120
C10A—C9A—C8A	120.7 (3)	C10B—C9B—C8B	120.4 (2)
С10А—С9А—Н9А	119.6	C10B—C9B—H9B	119.8
С8А—С9А—Н9А	119.6	C8B—C9B—H9B	119.8
C9A—C10A—C11A	120.3 (3)	C9B-C10B-C11B	120.3 (2)
C9A—C10A—H10A	119.8	C9B—C10B—H10B	119.8
C11A—C10A—H10A	119.8	C11B—C10B—H10B	119.8
C10A—C11A—C12A	119.6 (3)	C10B—C11B—C12B	120.0 (3)
C10A—C11A—H11A	120.2	C10B—C11B—H11B	120
C12A—C11A—H11A	120.2	C12B—C11B—H11B	120
C7A-C12A-C11A	120.0 (2)	C7B-C12B-C11B	119.4 (2)
C7A-C12A-C13A	120.8 (2)	C7B-C12B-C13B	120.5(2)
$C_{11A} = C_{12A} = C_{13A}$	119 2 (3)	C11B - C12B - C13B	120.0(2)
04A - C13A - C14A	119.2(3) 122 5 (3)	04B-C13B-C12B	120.0(2) 1193(2)
04A $-C13A$ $-C12A$	122.5(3) 1186(3)	O4B $C13B$ $C12B$ $O4B$ $C13B$ $C14B$	117.5(2)
C_{14A} C_{13A} C_{12A}	118.0(3)	C_{12B} C_{13B} C_{14B}	122.0(2) 118.0(2)
$C_{14A} = C_{13A} = C_{12A}$	110.9(2) 110.9(2)	C1B $C1AB$ $C5B$	110.0(2)
$C_{3A} = C_{14A} = C_{1A}$	119.9(2) 120.0(2)	C1P C14P C12P	119.5(2)
$C_{1A} = C_{14A} = C_{13A}$	120.0(2)	C_{1B} C_{14B} C_{13B} C_{5B} C_{14B} C_{12B}	120.0(2)
CIA = CI4A = CI3A	120.0(2)	C_{JB} C_{I4B} C_{IJB}	119.9 (2)
OIA = CI5A = U15A	108.4 (3)	OIB-CISB-CI6B	108.4 (2)
OIA—CISA—HISA	110	OIB-CISB-HISC	110
CI6A—CI5A—HI5A	110	CI6B—CI5B—HI5C	110
OIA—CI5A—HI5B	110	OIB—CI5B—HI5D	110
С16А—С15А—Н15В	110	C16B—C15B—H15D	110
H15A—C15A—H15B	108.4	H15C—C15B—H15D	108.4
C15A—C16A—H16A	109.5	C15B—C16B—H16D	109.5
C15A—C16A—H16B	109.5	C15B—C16B—H16E	109.5
H16A—C16A—H16B	109.5	H16D—C16B—H16E	109.5
C15A—C16A—H16C	109.5	C15B—C16B—H16F	109.5
H16A—C16A—H16C	109.5	H16D—C16B—H16F	109.5

H16B—C16A—H16C	109.5	H16E—C16B—H16F	109.5
O2A—C17A—C18A	107.4 (2)	O2B—C17B—C18B	107.5 (2)
O2A—C17A—H17A	110.2	O2B—C17B—H17C	110.2
C18A—C17A—H17A	110.2	C18B—C17B—H17C	110.2
O2A—C17A—H17B	110.2	O2B— $C17B$ — $H17D$	110.2
C18A—C17A—H17B	110.2	C18B—C17B—H17D	110.2
H17A—C17A—H17B	108.5	H17C-C17B-H17D	108.5
C17A—C18A—H18A	109.5	C17B-C18B-H18D	109.5
C17A - C18A - H18B	109.5	C17B— $C18B$ — $H18E$	109.5
H18A - C18A - H18B	109.5	H18D— $C18B$ — $H18E$	109.5
C17A - C18A - H18C	109.5	C17B-C18B-H18F	109.5
H18A - C18A - H18C	109.5	H18D— $C18B$ — $H18F$	109.5
H18B-C18A-H18C	109.5	H18F— $C18B$ — $H18F$	109.5
$C1A \rightarrow O1A \rightarrow C15A$	109.5 118 5 (2)	C1B - O1B - C15B	119.11 (19)
$C_{44} = 0.024 = C_{174}$	110.3(2) 119.1(2)	C4B - O2B - C17B	118.09(19)
C4A-02A-C1/A	119.1 (2)	C+D-02D-C17D	110.09 (19)
O1A—C1A—C2A—C3A	-178.3 (2)	O1B—C1B—C2B—C3B	175.1 (2)
C14A—C1A—C2A—C3A	1.0 (4)	C14B—C1B—C2B—C3B	-4.0 (4)
C1A—C2A—C3A—C4A	0.0 (4)	C1B—C2B—C3B—C4B	1.3 (4)
C2A—C3A—C4A—O2A	-179.9 (2)	C2B—C3B—C4B—O2B	-176.9 (2)
C2A—C3A—C4A—C5A	-0.2 (4)	C2B—C3B—C4B—C5B	2.0 (4)
O2A—C4A—C5A—C14A	179.0 (2)	O2B—C4B—C5B—C14B	176.5 (2)
C3A—C4A—C5A—C14A	-0.7 (4)	C3B—C4B—C5B—C14B	-2.4 (3)
O2A—C4A—C5A—C6A	-4.3 (4)	O2B—C4B—C5B—C6B	-3.0 (3)
C3A—C4A—C5A—C6A	176.0 (2)	C3B—C4B—C5B—C6B	178.1 (2)
C4A—C5A—C6A—O3A	-8.2 (4)	C4B—C5B—C6B—O3B	-17.1 (4)
C14A—C5A—C6A—O3A	168.5 (3)	C14B—C5B—C6B—O3B	163.4 (2)
C4A—C5A—C6A—C7A	175.7 (2)	C4B—C5B—C6B—C7B	163.8 (2)
C14A—C5A—C6A—C7A	-7.6 (4)	C14B—C5B—C6B—C7B	-15.7 (3)
O3A—C6A—C7A—C12A	-170.0 (3)	O3B—C6B—C7B—C12B	-161.9 (2)
C5A—C6A—C7A—C12A	6.3 (4)	C5B—C6B—C7B—C12B	17.2 (3)
O3A—C6A—C7A—C8A	8.4 (4)	O3B—C6B—C7B—C8B	15.1 (4)
C5A—C6A—C7A—C8A	-175.4 (2)	C5B—C6B—C7B—C8B	-165.8(2)
C12A—C7A—C8A—C9A	0.1 (4)	C12B—C7B—C8B—C9B	-0.9 (4)
C6A—C7A—C8A—C9A	-178.3 (2)	C6B—C7B—C8B—C9B	-177.9 (2)
C7A—C8A—C9A—C10A	0.1 (4)	C7B-C8B-C9B-C10B	1.6 (4)
C8A—C9A—C10A—C11A	-0.5 (5)	C8B-C9B-C10B-C11B	-0.7 (5)
C9A—C10A—C11A—C12A	0.6 (5)	C9B—C10B—C11B—C12B	-0.9 (4)
C8A—C7A—C12A—C11A	0.1 (4)	C8B-C7B-C12B-C11B	-0.6 (4)
C6A—C7A—C12A—C11A	178.4 (3)	C6B-C7B-C12B-C11B	176.3 (2)
C8A—C7A—C12A—C13A	-178.0 (3)	C8B—C7B—C12B—C13B	-179.2 (2)
C6A—C7A—C12A—C13A	0.3 (4)	C6B-C7B-C12B-C13B	-2.2 (3)
C10A—C11A—C12A—C7A	-0.4 (4)	C10B—C11B—C12B—C7B	1.5 (4)
C10A—C11A—C12A—C13A	177.7 (3)	C10B—C11B—C12B—C13B	-179.9 (3)
C7A—C12A—C13A—O4A	172.8 (4)	C7B—C12B—C13B—O4B	165.4 (2)
C11A—C12A—C13A—O4A	-5.4 (5)	C11B—C12B—C13B—O4B	-13.1 (4)
C7A—C12A—C13A—C14A	-5.6 (4)	C7B—C12B—C13B—C14B	-14.1 (3)
C11A—C12A—C13A—C14A	176.2 (3)	C11B—C12B—C13B—C14B	167.4 (2)

C4A—C5A—C14A—C1A C6A—C5A—C14A—C1A C4A—C5A—C14A—C13A C6A—C5A—C14A—C13A O1A—C1A—C14A—C5A C2A—C1A—C14A—C5A O1A—C1A—C14A—C5A O1A—C1A—C14A—C13A O4A—C13A—C14A—C13A O4A—C13A—C14A—C5A O4A—C13A—C14A—C5A O4A—C13A—C14A—C1A C12A—C13A—C14A—C1A C12A—C13A—C14A—C1A C12A—C13A—C14A—C1A C12A—C13A—O1A—C15A C14A—C1A—O1A—C15A C16A—C15A—O1A—C15A C16A—C15A—O1A—C17A C5A—C4A—O2A—C17A	$\begin{array}{c} 1.7 \ (4) \\ -175.0 \ (2) \\ 179.1 \ (3) \\ 2.4 \ (4) \\ 177.5 \ (2) \\ -1.8 \ (4) \\ 0.1 \ (4) \\ -179.2 \ (3) \\ -174.2 \ (4) \\ 4.1 \ (4) \\ 3.2 \ (5) \\ -178.4 \ (2) \\ 0.7 \ (4) \\ -178.6 \ (2) \\ 177.4 \ (2) \\ -4.4 \ (4) \\ 175.9 \ (2) \end{array}$	$\begin{array}{c} 01B-C1B-C14B-C5B\\ C2B-C1B-C14B-C5B\\ 01B-C1B-C14B-C13B\\ C2B-C1B-C14B-C13B\\ C2B-C1B-C14B-C13B\\ C4B-C5B-C14B-C1B\\ C6B-C5B-C14B-C13B\\ C4B-C5B-C14B-C13B\\ C6B-C5B-C14B-C13B\\ C6B-C5B-C14B-C13B\\ C12B-C13B-C14B-C1B\\ C12B-C13B-C14B-C5B\\ C12B-C13B-C14B-C15B\\ C14B-C1B-O1B-C15B\\ C16B-C15B-O1B-C1B\\ C3B-C4B-O2B-C17B\\ C5B-C4B-O2B-C17B\\ \end{array}$	$\begin{array}{c} -175.6 (2) \\ 3.5 (3) \\ 4.0 (3) \\ -176.8 (2) \\ -0.3 (3) \\ 179.2 (2) \\ 180.0 (2) \\ -0.5 (3) \\ 16.2 (4) \\ -164.3 (2) \\ -164.1 (2) \\ 15.4 (3) \\ -0.4 (4) \\ 178.7 (2) \\ -172.6 (2) \\ 5.2 (3) \\ -173.7 (2) \end{array}$
C5A—C4A—O2A—C17A	175.9 (2)	C5B—C4B—O2B—C17B	-173.7 (2)
C18A—C17A—O2A—C4A	179.5 (2)	C18B—C17B—O2B—C4B	175.7 (2)

Hydrogen-bond geometry (Å, °)

<i>D</i> —Н	H···A	$D \cdots A$	<i>D</i> —H… <i>A</i>
0.94	2.48	3.234 (3)	137
0.94	2.55	3.304 (4)	137
0.94	2.60	3.325 (3)	135
0.94	2.46	3.199 (4)	135
	<i>D</i> —H 0.94 0.94 0.94 0.94	D—H H···A 0.94 2.48 0.94 2.55 0.94 2.60 0.94 2.46	D—H H···A D···A 0.94 2.48 3.234 (3) 0.94 2.55 3.304 (4) 0.94 2.60 3.325 (3) 0.94 2.46 3.199 (4)

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