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Key indicators

Single-crystal X-ray study T = 150 K Mean $\sigma(\text{C-C}) = 0.006 \text{ Å}$ Disorder in main residue R factor = 0.024 wR factor = 0.054 Data-to-parameter ratio = 22.4

For details of how these key indicators were automatically derived from the article, see http://journals.iucr.org/e.

{2,6-Bis[(dimethylamino- κN)methyl]-4-{[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl}phenyl- κC^1 }(diiodine)iodidoplatinum(II) dichloromethane hemisolyate

In the title compound $[Pt(C_{17}H_{22}N_3O_4)I(I_2)] \cdot 0.5CH_2Cl_2$, the coordination environment of the Pt^{II} centre is distorted square-pyramidal and is defined by two neutral N atoms and an anionic C atom from the mono-anionic ligand, an iodide anion, and an additional η^1 -coordinated I_2 molecule. Intermolecular $C-H\cdots O$ contacts result in a dimeric structure. The asymmetric unit of (I) consists of two independent $[PtI(C_{17}H_{22}N_3O_4)(I_2)]$ molecules and a CH_2Cl_2 solvent molecula

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Comment

The study of the interaction of simple diatomic molecules with d^8 transition metals is crucial for a better understanding of the mechanism of oxidative addition and reductive elimination reactions. Therefore, we report here the crystal structure of the title compound, (I), as a product of the addition of I_2 to the pincer Pt^{II} complex 3,5-bis-[(dimethylamino)methyl]-4-[iodidoplatino(II)]benzoic acid N-hydroxysuccinimide ester.

$$I_2$$
 CH_2CI_2 / hexane
$$0.5CH_2CI_2$$

The asymmetric unit of (I) consists of two independent $[PtI(C_{17}H_{22}N_3O_4)(I_2)]$ molecules and a CH_2Cl_2 solvent molecule (Fig. 1). One molecule (containing Pt2) shows disorder

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metal-organic papers

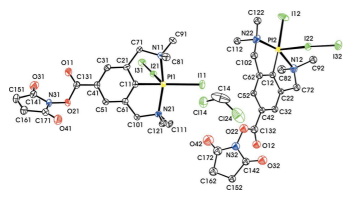


Figure 1
The asymmetric unit of (I), with the atomic numbering scheme. Displacement ellipsoids are drawn at the 50% probability level. H atoms have been omitted for clarity. Only one of the two disordered components is shown for molecule 2.

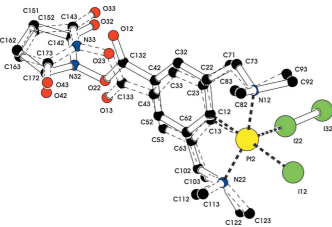


Figure 2
The disorder in the structure of the second molecule (Pt2). H atoms have been omitted for clarity. The minor disorder component is drawn with dashed bonds

about the molecular axis over all atomic positions, except for the heavy atoms and the N atoms bonded to Pt2 (Fig. 2). In the following, we discuss only molecule 1, containing Pt1. Geometric parameters for the Pt2 coordination environment are very similar to those for Pt1 and are given in Table 1.

The coordination environment of the PtII centre is defined the mono-anionic ligand 2,6-bis[(dimethylamino)methyl]-4-{[(2,5-dioxo-1-pyrrolidinyl)oxy]carbonyl}phenyl (C₁₇H₂₂N₃O₄), an iodide anion and an I₂ molecule. The ligand is coordinated in a tridentate manner to the metal via the anionic C atom [Pt1-C11 = 1.942 (4) Å] and the two neutral amine N atoms [Pt1-N11 = 2.104 (4) Å and Pt1-N12 =2.103 (4) Å]. The iodide anion is coordinated trans to atom C11 [C11-Pt1-I11 = 173.70 (12) $^{\circ}$], with a Pt1-I11 distance of 2.72985 (19) Å. The I_2 molecule is end-on η^1 -coordinated to the Pt^{II} centre [Pt1-I21 = 2.8260 (3) Å] and is part of a linear Pt1-I21-I23 arrangement [177.248 (14)°], with I21-I31 =2.8379 (4) Å. These distances compare well with those observed in similar Pt^{II} complexes, such as $[PtI(NCN)(\eta^1-I_2)]$ {NCN = 2,6-bis[(dimethylamino)methyl]phenyl, $C_{12}H_{19}N_2$ } reported by Gossage et al. (1999).

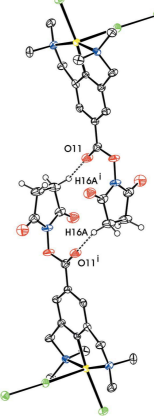


Figure 3 Hydrogen-bond interactions in (I). The $C-H\cdots O$ contacts are shown as dashed lines. [Symmetry code: (i) 1-x, -y, 1-z.]

The small bite angles of the chelate ligand [C11-Pt1-N11 = 82.09 (15) and C11-Pt1-N21 = 82.15 (15)°] result in a distorted square-pyramidal geometry of the central PtII atom. This can be quantified as 91.8% on the pathway from trigonal bipyramid to square pyramid (Holmes, 1984). The sum of the cis angles is 360°, although they deviate by up to 8° (for N21— Pt1-I11) from the ideal value of 90°. The largest deviation for the trans angles is 19° (N21-Pt1-N11) from the ideal value of 180°. A conformational analysis of ring puckering results in coefficients of 98.3° for the sine form of the PtC₃N11 chelate ring, and of 87.6° for the sine form of the PtC₃N21 chelate ring (Evans & Boeyens, 1989). Therefore, the two five-membered PtC₃N chelate rings are best described as twist conformations and are puckered in the same direction, with the N atoms mutually cis. The torsion angles Pt1-N11-C71-C21 = $26.1 (4)^{\circ}$ and $Pt1-N21-C101-C61 = -28.6 (4)^{\circ}$ result in a local non-crystallographic C_s symmetry.

The *N*-oxosuccinimide groups connect two molecules into a dimer structure *via* an intermolecular $C-H\cdots O$ contact, with $H16A\cdots O11^i=2.46\text{\AA}$ [symmetry code: (i) 1-x,-y,1-z] (Table 2 and Fig. 3).

Experimental

The reaction of 3,5-bis-[(dimethylamino)methyl]-4-[iodidoplatino(II)]benzoic acid, prepared as reported by Suijkerbuijk *et al.* (2002), with I_2 in CH_2Cl_2 -hexane (1:1) results in the title compound,

(I). Dark-red crystals suitable for X-ray data collection were obtained after recrystallization from CH₂Cl₂-hexane (1:1).

Crystal data

$[Pt(C_{17}H_{22}N_3O_4)I(I_2)]\cdot 0.5CH_2Cl_2$	$\gamma = 103.7589 \ (13)^{\circ}$
$M_r = 950.63$	$V = 2509.90 (7) \text{ Å}^3$
Triclinic, $P\overline{1}$	Z = 4
a = 8.7012 (1) Å	Mo $K\alpha$ radiation
b = 15.1862 (2) Å	$\mu = 9.41 \text{ mm}^{-1}$
c = 19.8391 (4) Å	T = 150 (2) K
$\alpha = 99.2137 \ (14)^{\circ}$	$0.42 \times 0.09 \times 0.06 \text{ mm}$
$\beta = 90.7250 \ (15)^{\circ}$	

Data collection

Nonius KappaCCD area-detector diffractometer Absorption correction: analytical (de Meulenaer & Tompa, 1965) $T_{\rm min}=0.14,\,T_{\rm max}=0.63$

35716 measured reflections 11455 independent reflections 8859 reflections with $I > 2\sigma(I)$ $R_{\rm int} = 0.036$

Refinement

 $\begin{array}{ll} R[F^2>2\sigma(F^2)]=0.024 & 162 \text{ restraints} \\ wR(F^2)=0.054 & \text{H-atom parameters constrained} \\ S=0.94 & \Delta\rho_{\max}=3.03 \text{ e Å}^{-3} \\ 11455 \text{ reflections} & \Delta\rho_{\min}=-3.35 \text{ e Å}^{-3} \end{array}$

 Table 1

 Selected geometric parameters (\mathring{A} , °).

Pt1-I21 2.8260 (3) Pt2-I22 2.8198 (3) I21-I31 2.8379 (4) I22-I32 2.8472 (4) Pt2-C13 1.908 (9) 1.22-I32 2.8472 (4) C11-Pt1-N21 82.15 (15) C13-Pt2-N12 85.7 (3) C11-Pt1-N11 161.06 (14) N22-Pt2-N12 160.14 (16) C11-Pt1-I11 173.70 (12) C13-Pt2-I12 173.3 (4) N21-Pt1-I11 98.26 (10) C12-Pt2-I12 172.6 (4) N11-Pt1-I11 96.26 (10) N22-Pt2-I12 97.20 (12) C11-Pt1-I21 84.54 (12) N12-Pt2-I12 97.75 (11) N21-Pt1-I21 94.22 (10) C13-Pt2-I22 83.7 (5) N11-Pt1-I21 94.68 (10) C12-Pt2-I22 85.4 (41) N11-Pt1-I21 101.675 (11) N22-Pt2-I22 94.64 (11) Pt1-I21-I31 177.248 (14) N12-Pt2-I22 95.10 (10) C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt2-I22-I32 177.533 (15)				
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Pt1-I11 2.6862 (3) Pt2-I12 2.6827 (4) Pt1-I21 2.8260 (3) Pt2-I22 2.8198 (3) I21-I31 2.8379 (4) I22-I32 2.8472 (4) Pt2-C13 1.908 (9) 2.8472 (4) C11-Pt1-N21 82.15 (15) C13-Pt2-N12 85.7 (3) C11-Pt1-N11 161.06 (14) N22-Pt2-N12 79.3 (3) N21-Pt1-N11 161.06 (14) N22-Pt2-N12 160.14 (16) C11-Pt1-I11 173.70 (12) C13-Pt2-I12 173.3 (4) N21-Pt1-I11 98.26 (10) C12-Pt2-I12 172.6 (4) N11-Pt1-I11 96.26 (10) N22-Pt2-I12 97.20 (12) C11-Pt1-I21 84.54 (12) N12-Pt2-I12 97.55 (11) N21-Pt1-I21 94.22 (10) C13-Pt2-I22 83.7 (5) N11-Pt1-I21 94.68 (10) C12-Pt2-I22 85.4 (41) N11-Pt1-I21 101.675 (11) N22-Pt2-I22 94.64 (11) Pt1-I21-I31 177.248 (14) N12-Pt2-I22 95.10 (10) C13-Pt2-C12 6.5 (4) I12-Pt2-I22 101.	Pt1-N21	2.103 (4)	Pt2-N22	2.098 (4)
Pt1-I21 2.8260 (3) Pt2-I22 2.8198 (3) I21-I31 2.8379 (4) I22-I32 2.8472 (4) Pt2-C13 1.908 (9) 1.22-I32 2.8472 (4) C11-Pt1-N21 82.15 (15) C13-Pt2-N12 85.7 (3) C11-Pt1-N11 82.09 (15) C12-Pt2-N12 79.3 (3) N21-Pt1-N11 161.06 (14) N22-Pt2-N12 160.14 (16) C11-Pt1-I11 173.70 (12) C13-Pt2-I12 173.3 (4) N21-Pt1-I11 98.26 (10) C12-Pt2-I12 172.6 (4) N11-Pt1-I11 96.26 (10) N22-Pt2-I12 97.20 (12) C11-Pt1-I21 84.54 (12) N12-Pt2-I12 97.55 (11) N21-Pt1-I21 94.68 (10) C13-Pt2-I22 83.7 (5) N11-Pt1-I21 94.68 (10) C12-Pt2-I22 85.4 (4) I11-Pt1-I21 101.675 (11) N22-Pt2-I22 95.10 (10) C13-Pt2-C12 6.5 (4) I12-Pt2-I22 95.10 (10) C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt2-I	Pt1-N11	2.104 (4)	Pt2-N12	2.103 (4)
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Pt2—C13 1.908 (9) C11—Pt1—N21 82.15 (15) C13—Pt2—N12 85.7 (3) C11—Pt1—N11 82.09 (15) C12—Pt2—N12 79.3 (3) N21—Pt1—N11 161.06 (14) N22—Pt2—N12 160.14 (16) C11—Pt1—I11 173.70 (12) C13—Pt2—I12 173.3 (4) N21—Pt1—I11 98.26 (10) C12—Pt2—I12 172.6 (4) N11—Pt1—I11 96.26 (10) N22—Pt2—I12 97.20 (12) C11—Pt1—I21 84.54 (12) N12—Pt2—I12 97.75 (11) N21—Pt1—I21 94.22 (10) C13—Pt2—I22 83.7 (5) N11—Pt1—I21 94.68 (10) C12—Pt2—I22 85.4 (4) II1—Pt1—I21 101.675 (11) N22—Pt2—I22 94.64 (11) Pt1—I21—I31 177.248 (14) N12—Pt2—I22 95.10 (10) C13—Pt2—C12 6.5 (4) I12—Pt2—I22 101.569 (11) C13—Pt2—N22 78.2 (3) Pt2—I22—I32 177.533 (15) C12—Pt2—N22 84.3 (3) Pt2—I22—C102—C62 -28.4 (7) Pt1—N11—C71—C21 26.1 (4) Pt2—N22—C102—C62 <td< td=""><td>Pt1-I21</td><td>2.8260 (3)</td><td>Pt2-I22</td><td>2.8198 (3)</td></td<>	Pt1-I21	2.8260 (3)	Pt2-I22	2.8198 (3)
C11-Pt1-N21 82.15 (15) C13-Pt2-N12 85.7 (3) C11-Pt1-N11 82.09 (15) C12-Pt2-N12 79.3 (3) N21-Pt1-N11 161.06 (14) N22-Pt2-N12 160.14 (16) C11-Pt1-I11 173.70 (12) C13-Pt2-I12 173.3 (4) N21-Pt1-I11 98.26 (10) C12-Pt2-I12 172.6 (4) N11-Pt1-I11 96.26 (10) N22-Pt2-I12 97.20 (12) C11-Pt1-I21 84.54 (12) N12-Pt2-I12 97.75 (11) N21-Pt1-I21 94.22 (10) C13-Pt2-I22 83.7 (5) N11-Pt1-I21 94.68 (10) C12-Pt2-I22 85.4 (4) I11-Pt1-I21 101.675 (11) N22-Pt2-I22 94.64 (11) Pt1-I21-I31 177.248 (14) N12-Pt2-I22 95.10 (10) C13-Pt2-C12 6.5 (4) N12-Pt2-I22 101.569 (11) C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	I21-I31	2.8379 (4)	I22-I32	2.8472 (4)
C11-Pt1-N11 82.09 (15) C12-Pt2-N12 79.3 (3) N21-Pt1-N11 161.06 (14) N22-Pt2-N12 160.14 (16) C11-Pt1-I11 173.70 (12) C13-Pt2-I12 173.3 (4) N21-Pt1-I11 98.26 (10) C12-Pt2-I12 172.6 (4) N11-Pt1-I11 96.26 (10) N22-Pt2-I12 97.20 (12) C11-Pt1-I21 84.54 (12) N12-Pt2-I12 97.75 (11) N21-Pt1-I21 94.22 (10) C13-Pt2-I22 83.7 (5) N11-Pt1-I21 94.68 (10) C12-Pt2-I22 85.4 (4) I11-Pt1-I21 101.675 (11) N22-Pt2-I22 94.64 (11) Pt1-I21-I31 177.248 (14) N12-Pt2-I22 95.10 (10) C13-Pt2-C12 6.5 (4) I12-Pt2-I22 101.569 (11) C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	Pt2-C13	1.908 (9)		
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N11-Pt1-I11 96.26 (10) N22-Pt2-I12 97.20 (12) C11-Pt1-I21 84.54 (12) N12-Pt2-I12 97.75 (11) N21-Pt1-I21 94.22 (10) C13-Pt2-I22 83.7 (5) N11-Pt1-I21 94.68 (10) C12-Pt2-I22 85.4 (41) I11-Pt1-I21 101.675 (11) N22-Pt2-I22 94.64 (11) Pt1-I21-I31 177.248 (14) N12-Pt2-I22 95.10 (10) C13-Pt2-C12 6.5 (4) I12-Pt2-I22 101.569 (11) C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	C11-Pt1-I11	173.70 (12)	C13-Pt2-I12	173.3 (4)
C11-Pt1-I21 84.54 (12) N12-Pt2-I12 97.75 (11) N21-Pt1-I21 94.22 (10) C13-Pt2-I22 83.7 (5) N11-Pt1-I21 94.68 (10) C12-Pt2-I22 85.4 (4) I11-Pt1-I21 101.675 (11) N22-Pt2-I22 94.64 (11) Pt1-I21-I31 177.248 (14) N12-Pt2-I22 95.10 (10) C13-Pt2-C12 6.5 (4) I12-Pt2-I22 101.569 (11) C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	N21-Pt1-I11	98.26 (10)	C12-Pt2-I12	172.6 (4)
N21-Pt1-I21 94.22 (10) C13-Pt2-I22 83.7 (5) N11-Pt1-I21 94.68 (10) C12-Pt2-I22 85.4 (4) II1-Pt1-I21 101.675 (11) N22-Pt2-I22 94.64 (11) Pt1-I21-I31 177.248 (14) N12-Pt2-I22 95.10 (10) C13-Pt2-C12 6.5 (4) I12-Pt2-I22 101.569 (11) C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	N11-Pt1-I11	96.26 (10)	N22-Pt2-I12	97.20 (12)
N11-Pt1-I21 94.68 (10) C12-Pt2-I22 85.4 (4) I11-Pt1-I21 101.675 (11) N22-Pt2-I22 94.64 (11) Pt1-I21-I31 177.248 (14) N12-Pt2-I22 95.10 (10) C13-Pt2-C12 6.5 (4) I12-Pt2-I22 101.569 (11) C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	C11-Pt1-I21	84.54 (12)	N12-Pt2-I12	97.75 (11)
I11-Pt1-I21 101.675 (11) N22-Pt2-I22 94.64 (11) Pt1-I21-I31 177.248 (14) N12-Pt2-I22 95.10 (10) C13-Pt2-C12 6.5 (4) I12-Pt2-I22 101.569 (11) C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	N21-Pt1-I21	94.22 (10)	C13-Pt2-I22	83.7 (5)
Pt1-I21-I31 177.248 (14) N12-Pt2-I22 95.10 (10) C13-Pt2-C12 6.5 (4) I12-Pt2-I22 101.569 (11) C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	N11-Pt1-I21	94.68 (10)	C12-Pt2-I22	85.4 (4)
C13-Pt2-C12 6.5 (4) I12-Pt2-I22 101.569 (11) C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	I11-Pt1-I21	101.675 (11)	N22-Pt2-I22	94.64 (11)
C13-Pt2-N22 78.2 (3) Pt2-I22-I32 177.533 (15) C12-Pt2-N22 84.3 (3) Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	Pt1-I21-I31	177.248 (14)	N12-Pt2-I22	95.10 (10)
C12-Pt2-N22 84.3 (3) Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	C13-Pt2-C12	6.5 (4)	I12-Pt2-I22	101.569 (11)
Pt1-N11-C71-C21 26.1 (4) Pt2-N22-C102-C62 -28.4 (7) Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	C13-Pt2-N22	78.2 (3)	Pt2-I22-I32	177.533 (15)
Pt1-N21-C101-C61 -28.6 (4) Pt2-N12-C73-C23 20.5 (11)	C12-Pt2-N22	84.3 (3)		
	Pt1-N11-C71-C21	26.1 (4)	Pt2-N22-C102-C62	-28.4(7)
Pt2-N12-C72-C22 25.9 (9) Pt2-N22-C103-C63 -35.1 (9)	Pt1-N21-C101-C61	-28.6(4)	Pt2-N12-C73-C23	20.5 (11)
	Pt2-N12-C72-C22	25.9 (9)	Pt2-N22-C103-C63	-35.1 (9)

Table 2 Hydrogen-bond geometry (Å, °).

D $ H$ $\cdots A$	D-H	$H \cdot \cdot \cdot A$	$D \cdot \cdot \cdot A$	$D-\mathrm{H}\cdots A$
C161—H16A···O11i	0.99	2.46	3.296 (7)	142

Symmetry code: (i) 1 - x, -y, 1 - z.

All H atoms were introduced in geometrically idealized positions, with C–H = 0.95–0.99 Å and refined using a riding model, with $U_{\rm iso}({\rm H})=1.2U_{\rm eq}({\rm C})$ for CH and CH $_{\rm 2}$ H atoms, or $1.5U_{\rm eq}({\rm C})$ for methyl H atoms. Disordered C, N and O atoms were refined isotropically. The highest peak and deepest hole in the residual electron density are both 0.8 Å from atom I12. All C and O atoms and atom N32 in the second molecule (Pt2) are disordered about the long axis of the molecule. The occupancy factors for these positions were refined to 0.566 (6) and 0.434 (6). Atoms C41/C131/O11/O21/N31, C42/C132/O12/O22/N32 and C42/C133/O13/O23/N33 were restrained to lie on respective common planes.

Data collection: *COLLECT* (Nonius, 1999); cell refinement: *DIRAX* (Duisenberg, 1992); data reduction: *EVALCCD* (Duisenberg *et al.*, 2003) and *SORTAV* (Blessing, 1987); program(s) used to solve structure: *DIRDIF97* (Beurskens *et al.*, 1997); program(s) used to refine structure: *SHELXL97* (Sheldrick, 1997); molecular graphics: *PLATON* (Spek, 2003); software used to prepare material for publication: manual editing of the *SHELXL97* output.

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