V = 1441.91 (15) Å<sup>3</sup>

 $0.61 \times 0.18 \times 0.08 \text{ mm}$ 

Diffraction, 2010)

 $T_{\rm min} = 0.780, \; T_{\rm max} = 1$ 

7277 measured reflections 2685 independent reflections

2021 reflections with  $I > 2\sigma(I)$ 

Mo  $K\alpha$  radiation

 $\mu = 0.22 \text{ mm}^{-1}$ 

T = 298 K

 $R_{\rm int} = 0.031$ 

Z = 2

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# 3,3'-Dibenzoyl-1,1'-dibenzyl-1,1'-(ethane-1,2-diyl)dithiourea

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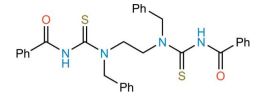
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Key indicators: single-crystal X-ray study; T = 298 K; mean  $\sigma$ (C–C) = 0.004 Å; disorder in main residue; R factor = 0.050; wR factor = 0.123; data-to-parameter ratio = 11.6.

In the title compound,  $C_{32}H_{30}N_4O_2S_2$ , the carbonyl and thiocarbonyl groups are found in a rare synclinal conformation, with an S–C···C–O pseudo-torsion angle of 62.6 (2)°. The molecule has  $C_i = S_2$  point-group symmetry with a crystallographic center of inversion located in the middle of the ethylene bridge. One of the symmetry-independent phenyl rings is disordered over two orientations, with a siteoccupation ratio of 70:30. The distances between the centroids of the nearest phenyl rings are equal to one of the lattice constants [a = 4.7767 (2) Å], so stacking interactions are extremely weak. Molecules are joined by bifurcated hydrogen bonds (N-H···O and N-H···S), forming a ladder-like arrangement along [100]. van der Waals forces combine these ladders into a three-dimensional structure. The dependency between the  $S \cdots O$  distance and the improper  $S = C \cdots C = O$ torsion angle based on 739 structures containing the CC(=O)NC(=S)N moiety is discussed.

#### **Related literature**

For structures of bis(N-benzoylthioureas) derived from aliphatic diamines, see: Ding et al. (2008); Dong et al. (2007); Sow et al. (2009). For those derived from o-cyclohexanediamine, see: Jumal et al. (2011). For those derived from aromatic diamines, see: Cao et al. (2007); Li et al. (2009); Thiam et al. (2008); Woei Hung & Kassim (2010); Yamin & Osman (2011). For other acyl derivaties obtained from o- and p-phenylenediamine (also solvates), see: Dong, Yan et al. (2008); Dong, Yang et al. (2008); Du & Du (2008); Du et al. (2008). For 1-benzoyl-3-phenylurea, see: Okuniewski et al. (2010). For the synthetic procedure, see: Douglass & Dains (1934). For a review on N-aroylthioureas, see: Aly et al. (2007). For a description of the Cambridge Structural Database, see: Allen (2002).



#### **Experimental**

Crystal data C32H30N4O2S2  $M_{\rm m} = 566.72$ Monoclinic,  $P2_1/c$ a = 4.7767 (2) Å b = 25.1653 (16) Å c = 11.9998 (8) Å  $\beta = 91.585(5)^{\circ}$ 

#### Data collection

Oxford Diffraction Xcalibur diffractometer with a Sapphire2 (large Be window) detector Absorption correction: multi-scan (CrysAlis PRO; Oxford

#### Refinement

| $R[F^2 > 2\sigma(F^2)] = 0.05$ | H atoms treated by a mixture of                           |
|--------------------------------|---|
| $wR(F^2) = 0.123$              | independent and constrained                               |
| S = 1.04                       | refinement  |
| 2685 reflections               | $\Delta \rho_{\rm max} = 0.30 \text{ e } \text{\AA}^{-3}$ |
| 231 parameters                 | $\Delta \rho_{\rm min} = -0.15 \text{ e} \text{ Å}^{-3}$  |
| 163 restraints                 |   |

#### Table 1

Hydrogen-bond geometry (Å, °).

| $D - H \cdot \cdot \cdot A$       | D-H      | $H \cdot \cdot \cdot A$ | $D \cdots A$ | $D - \mathbf{H} \cdots A$ |
|-----------------------------------|----------|-------------------------|--------------|---------------------------|
| $N1 - H1N \cdots O1^i$            | 0.86 (1) | 2.30(1)                 | 3.073 (2)    | 150 (2)                   |
| $N1 - H1N \cdot \cdot \cdot S1^i$ | 0.86(1)  | 2.98 (2)                | 3.647 (2)    | 136 (2)                   |

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

Data collection: CrysAlis PRO (Oxford Diffraction, 2010); cell refinement: CrysAlis PRO; data reduction: CrysAlis PRO; program(s) used to solve structure: SHELXS97 (Sheldrick, 2008); program(s) used to refine structure: SHELXL97 (Sheldrick, 2008); molecular graphics: OLEX2 (Dolomanov et al., 2009); software used to prepare material for publication: WinGX (Farrugia, 1999) and PLATON (Spek, 2009).

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Supplementary data and figures for this paper are available from the IUCr electronic archives (Reference: IM2351).

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# supporting information

Acta Cryst. (2012). E68, o619-o620 [doi:10.1107/S1600536812002954]

# 3,3'-Dibenzoyl-1,1'-dibenzyl-1,1'-(ethane-1,2-diyl)dithiourea

## Andrzej Okuniewski, Jaroslaw Chojnacki and Barbara Becker

#### S1. Comment

Substituted *N*-acylthioureas are the subject of extensive research because of their biological activity, metal coordination ability and hydrogen bond formation (Aly *et al.*, 2007). Bis(*N*-acylthioureas) are a reatively less studied group with respect to their mono-analogues (Yamin & Osman, 2011).

The title compound,  $[PhCONHCSN(CH_2Ph)CH_2]_2$  (Fig.1), has the inversion center located in the middle of the ethylene bridge, so only half of the molecule is symetrically independent. There is no intramolecular N—H···O hydrogen bond that is commonly present in substituted *N*-acylthioureas and ureas (Okuniewski *et al.*, 2010), because the hydrogen atom on N2 is substituted by a benzyl group. The only specific interactions are weak bifurcative N—H···O and N—H···S intermolecular hydrogen bonds joining molecules into one-dimensional ladders along [100] (Fig. 2). The threedimensional structure is only stabilized by van der Waals forces (Fig. 3).

Three main geometries of *N*-acylthioureas based on the S···O distance ( $d_{SO}$ ) and the S=C···C=O improper torsion angle ( $\varphi_{SCCO}$ ) can be distinguished: synperiplanar type (i) and (iii) with  $|\varphi_{SCCO}| \approx 0^{\circ}$  as well as antiperiplanar type (ii) with |  $\varphi_{SCCO}| \approx 180^{\circ}$  (Fig. 4). The S···O distance in type (i) is about 3 Å while in type (iii) is about 5 Å. Transition between type (i) and (ii) is smooth and is accomplished by rotation about the thioamide bond. Theoretical relation between  $d_{SO}$  and  $\varphi_{SCCO}$  assuming constant bond lengths and valence angles can be expressed as (see: solid line in Fig. 4):

 $d_{\rm SO} = (A \cos \varphi_{\rm SCCO} + B)^{0.5}$ 

where A and B are calculated as:

 $A = 2 d_{\rm SC} \sin \alpha_{\rm SCN} \left[ d_{\rm CO} \sin(\alpha_{\rm NCO} + \alpha_{\rm CNC}) - d_{\rm NC} \sin \alpha_{\rm CNC} \right] \simeq -6.4657 \text{ Å}^2$ 

 $B = [-d_{\rm CN} + d_{\rm CN} \cos \alpha_{\rm CNC} - d_{\rm CO} \cos(\alpha_{\rm NCO} + \alpha_{\rm CNC}) + d_{\rm SC} \cos \alpha_{\rm SCN}]^2 + [-d_{\rm SC} \sin \alpha_{\rm SCN}]^2 + [d_{\rm NC} \sin \alpha_{\rm CNC} - d_{\rm CO} \sin(\alpha_{\rm NCO} + \alpha_{\rm CNC})]^2 \\ \approx 13.666 \text{ Å}^2$ 

Numerical values of bond lengths and angles are the average ones calculated in Vista program on the basis of 739 structures (980 values) containing CC(=O)NC(=S)N moiety found in CSD 5.32 (Allen, 2002)

Type (ii) is generally more stable than type (i) due to the formation of the intramolecular N—H···O hydrogen bond. When there is no suitable hydrogen atom to form hydrogen bonds (*N*,*N*-disubstituted derivatives) the anticlinal geometry ( $|\varphi_{SCCO}| \approx 120^{\circ}$ ) is preferred. Only 58 out of 980 points in Fig. 4 represent type (i) with  $d_{SO} < 4$  Å and  $|\varphi_{SCCO}| < 90^{\circ}$ . In this type the S···O distance is slightly greater than theoretical value due to sulfur-oxygen repulsion. Molecules of type (iii) contain covalent six-membered rings (see: structure (iii) in Fig. 4) preventing any rotation, so there is no possibility to transform this type to any other.

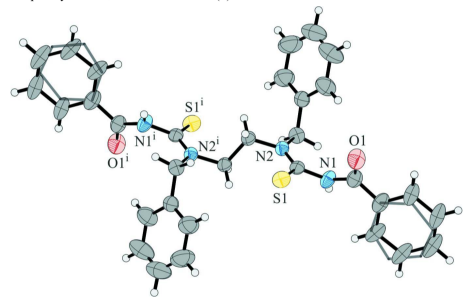
Geometric parameters of the title compound's molecule place it near type (i) – rare synclinal conformation with  $|\varphi_{SCCO}| = 62.60 (18)^{\circ}$  (see: cross mark in Fig. 4). Large substituents on N2 atom cause type (ii) to be geometrically unfavourable.

## **S2. Experimental**

Synthesis was performed according to Douglass & Dains (1934): 2.50 g (33 mmol) of ammonium thiocyanate and 20 ml of acetone were placed in a two-necked flask. Through a dropping funnel 3.49 ml (30 mmol) of benzoyl chloride in 20 ml of acetone was added with stirring. After addition was completed the mixture was refluxed for additional 15 min and then 3.54 ml (15 mmol) of *N*,*N*'-dibenzylethane-1,2-diamine in 20 ml of acetone was added through the dropping funnel. The mixture was carefully poured to the 500 ml of water with stirring. The resulting precipitate was filtered on a Büchner funnel. The crude product was recrystallized from acetone. Colorless single crystals suitable for X-ray diffraction analysis were isolated with 72% yield. Melting point:  $165 (1)^{\circ}C$ .

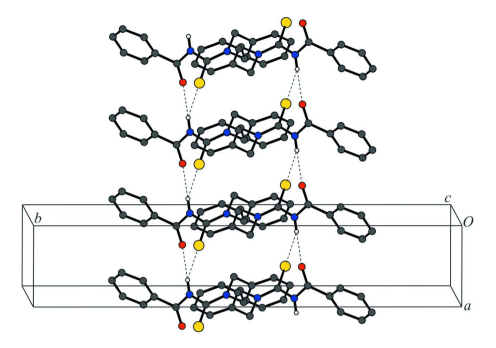
#### **S3. Refinement**

All C-bonded hydrogen atoms were placed in calculated positions (aromatic:  $d_{CH} = 0.97$  Å, methylene:  $d_{CH} = 0.93$  Å) and were treated as riding on their parent atoms with  $U_{iso}$  (H) = 1.2  $U_{eq}$ (C). H1N atom was located from difference Fourier map and refined isotropically with  $d_{NH}$  restrained to 0.88 (1) Å.



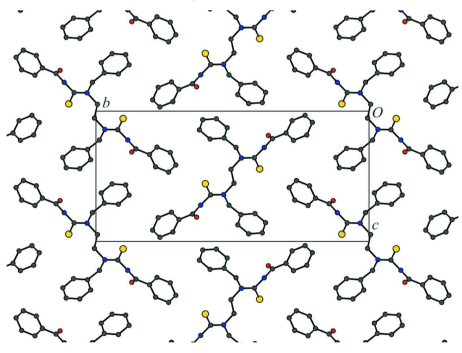
## Figure 1

The molecular structure of the title compound with displacement ellipsoids drawn at the 50% probability level. Symmetry code: (i) -x + 2, -y + 1, -z + 1.



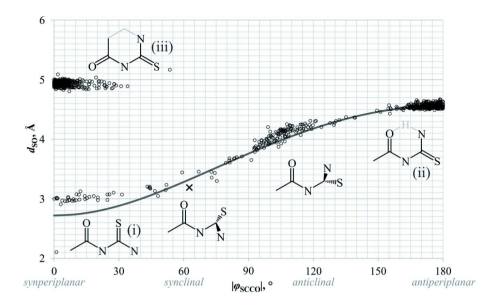
# Figure 2

Ladder-like structural motif found in the structure of the title compound. Hydrogen bonds are shown as dashed lines. Cbonded hydrogen atoms have been ommited for clarity.



# Figure 3

Tiling of ladders seen along [100] direction. Hydrogen atoms have been ommited for clarity.



# Figure 4

Plot of S…O distance ( $d_{SO}$ ) against absolute value of S=C…C=O torsion angle ( $|\varphi_{SCCO}|$ ) for 739 structures containing CC(=O)NC(=S)N fragment found in CSD 5.32 (Allen, 2002). Title compound is marked with cross. Solid line represents theoretical relation (see: comment).

## 3-benzoyl-1-benzyl-1-(2- {benzyl[(phenylformamido)methanethioyl]amino}ethyl)thiourea

| Crystal data  |   |
|---|---|
| $C_{32}H_{30}N_4O_2S_2$                             | F(000) = 596  |
| $M_r = 566.72$                                      | $D_{\rm x} = 1.305 {\rm ~Mg} {\rm ~m}^{-3}$                         |
| Monoclinic, $P2_1/c$                                | Melting point: 438(1) K   |
| Hall symbol: -P 2ybc                                | Mo <i>K</i> $\alpha$ radiation, $\lambda = 0.71073$ Å               |
| a = 4.7767 (2)  Å                                   | Cell parameters from 2955 reflections                               |
| b = 25.1653 (16)  Å                                 | $\theta = 2.3 - 28.7^{\circ}$                                       |
| c = 11.9998 (8) Å                                   | $\mu = 0.22 \mathrm{~mm^{-1}}$                                      |
| $\beta = 91.585(5)^{\circ}$                         | T = 298  K  |
| $V = 1441.91 (15) Å^3$                              | Needle, colourless  |
| Z = 2   | $0.61\times0.18\times0.08~mm$                                       |
| Data collection                                     |   |
| Oxford Diffraction Xcalibur Sapphire2 (large        | 7277 measured reflections   |
| Be window)  | 2685 independent reflections  |
| diffractometer                                      | 2021 reflections with $I > 2\sigma(I)$                              |
| Graphite monochromator                              | $R_{\rm int} = 0.031$   |
| Detector resolution: 8.1883 pixels mm <sup>-1</sup> | $\theta_{\rm max} = 25.5^{\circ}, \ \theta_{\rm min} = 2.4^{\circ}$ |
| $\omega$ scans                                      | $h = -5 \rightarrow 5$  |
| Absorption correction: multi-scan                   | $k = -26 \rightarrow 30$  |
| (CrysAlis PRO; Oxford Diffraction, 2010)            | $l = -14 \rightarrow 14$  |
| $T_{\min} = 0.780, \ T_{\max} = 1$                  |   |

Refinement

| 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                                 |
| $w = 1/[\sigma^2(F_o^2) + (0.0551P)^2 + 0.2801P]$          |
| where $P = (F_o^2 + 2F_c^2)/3$                             |
| $(\Delta/\sigma)_{\rm max} = 0.016$                        |
| $\Delta \rho_{\rm max} = 0.30 \text{ e } \text{\AA}^{-3}$  |
| $\Delta \rho_{\rm min} = -0.15 \text{ e } \text{\AA}^{-3}$ |
|  |

#### Special details

Experimental. CrysAlisPro, Oxford Diffraction Ltd., Version 1.171.33.66 (Oxford Diffraction, 2010) Empirical

absorption correction using spherical harmonics, implemented in SCALE3 ABSPACK scaling algorithm. **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 of  $F^2$  against ALL reflections. The weighted *R*-factor *wR* and goodness of fit *S* are based on  $F^2$ , conventional *R*-factors *R* are based on *F*, with *F* set to zero for negative  $F^2$ . The threshold expression of  $F^2 > \sigma(F^2)$  is used only for calculating *R*-factors(gt) *etc.* and is not relevant to the choice of reflections for refinement. *R*-factors based on  $F^2$ are statistically about twice as large as those based on *F*, and *R*- factors based on ALL data will be even larger.

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

|      | x          | У           | Ζ            | $U_{ m iso}$ */ $U_{ m eq}$ | Occ. (<1) |
|------|------------|-------------|--------------|-----------------------------|-----------|
| C1   | 1.1371 (4) | 0.58001 (8) | 0.64272 (17) | 0.0356 (5)                  |           |
| C3   | 1.0855 (4) | 0.49446 (8) | 0.55324 (16) | 0.0377 (5)                  |           |
| H3A  | 1.0468     | 0.4587      | 0.5788       | 0.045*                      |           |
| H3B  | 1.2835     | 0.4967      | 0.5379       | 0.045*                      |           |
| C10  | 1.2067 (5) | 0.64605 (9) | 0.7895 (2)   | 0.0458 (5)                  |           |
| C11  | 1.0914 (5) | 0.69614 (9) | 0.8323 (2)   | 0.0528 (6)                  |           |
| C12A | 0.882 (3)  | 0.7252 (6)  | 0.7782 (14)  | 0.065 (3)                   | 0.71 (3)  |
| H12A | 0.7981     | 0.713       | 0.7121       | 0.078*                      | 0.71 (3)  |
| C13A | 0.800(2)   | 0.7731 (4)  | 0.8256 (11)  | 0.077 (3)                   | 0.71 (3)  |
| H13A | 0.6566     | 0.7927      | 0.7911       | 0.092*                      | 0.71 (3)  |
| C14A | 0.924 (2)  | 0.7922 (3)  | 0.9218 (13)  | 0.078 (3)                   | 0.71 (3)  |
| H14A | 0.8664     | 0.8244      | 0.9513       | 0.093*                      | 0.71 (3)  |
| C15A | 1.138 (3)  | 0.7634 (4)  | 0.9756 (10)  | 0.078 (3)                   | 0.71 (3)  |
| H15A | 1.2214     | 0.7762      | 1.0413       | 0.093*                      | 0.71 (3)  |
| C16A | 1.225 (3)  | 0.7153 (5)  | 0.9299 (9)   | 0.065 (2)                   | 0.71 (3)  |
| H16A | 1.3702     | 0.6961      | 0.9638       | 0.077*                      | 0.71 (3)  |
| C12B | 0.873 (7)  | 0.7181 (15) | 0.769 (3)    | 0.075 (7)                   | 0.29 (3)  |
| H12B | 0.8008     | 0.6968      | 0.7118       | 0.091*                      | 0.29 (3)  |
| C13B | 0.747 (7)  | 0.7674 (12) | 0.780 (3)    | 0.097 (6)                   | 0.29 (3)  |
| H13B | 0.6192     | 0.7814      | 0.7281       | 0.116*                      | 0.29 (3)  |
| C14B | 0.831 (7)  | 0.7934 (10) | 0.875 (2)    | 0.083 (6)                   | 0.29 (3)  |
| H14B | 0.7374     | 0.8243      | 0.8941       | 0.099*                      | 0.29 (3)  |

| C15B | 1.048 (7)    | 0.7760 (10)  | 0.943 (3)    | 0.080 (7)   | 0.29 (3) |
|------|--------------|--------------|--------------|-------------|----------|
| H15B | 1.1111       | 0.7964       | 1.0035       | 0.095*      | 0.29 (3) |
| C16B | 1.168 (7)    | 0.7275 (11)  | 0.920 (2)    | 0.075 (7)   | 0.29 (3) |
| H16B | 1.3107       | 0.7155       | 0.9683       | 0.09*       | 0.29 (3) |
| C20  | 0.8164 (4)   | 0.51283 (9)  | 0.72178 (18) | 0.0410 (5)  |          |
| H20A | 0.6511       | 0.4995       | 0.6822       | 0.049*      |          |
| H20B | 0.7592       | 0.5421       | 0.7687       | 0.049*      |          |
| C21  | 0.9395 (4)   | 0.46935 (9)  | 0.79385 (17) | 0.0423 (5)  |          |
| C22  | 1.1471 (6)   | 0.48056 (12) | 0.8727 (2)   | 0.0678 (8)  |          |
| H22  | 1.2159       | 0.515        | 0.8798       | 0.081*      |          |
| C23  | 1.2529 (6)   | 0.44077 (16) | 0.9412 (3)   | 0.0881 (10) |          |
| H23  | 1.395        | 0.4487       | 0.9931       | 0.106*      |          |
| C24  | 1.1543 (7)   | 0.39089 (16) | 0.9341 (3)   | 0.0861 (11) |          |
| H24  | 1.2225       | 0.3648       | 0.9826       | 0.103*      |          |
| C25  | 0.9545 (8)   | 0.37876 (13) | 0.8557 (3)   | 0.0920 (11) |          |
| H25  | 0.8906       | 0.344        | 0.8487       | 0.11*       |          |
| C26  | 0.8446 (6)   | 0.41794 (11) | 0.7858 (2)   | 0.0704 (8)  |          |
| H26  | 0.7056       | 0.4093       | 0.733        | 0.085*      |          |
| N1   | 1.0341 (4)   | 0.61630 (7)  | 0.72100 (16) | 0.0442 (5)  |          |
| N2   | 1.0180 (3)   | 0.53247 (7)  | 0.64038 (13) | 0.0345 (4)  |          |
| O1   | 1.4457 (3)   | 0.63131 (7)  | 0.81329 (15) | 0.0619 (5)  |          |
| S1   | 1.38677 (11) | 0.59942 (2)  | 0.55656 (5)  | 0.0483 (2)  |          |
| H1N  | 0.855 (2)    | 0.6203 (9)   | 0.7200 (19)  | 0.053 (7)*  |          |
|      |              |              |              |             |          |

Atomic displacement parameters  $(Å^2)$ 

|      | $U^{11}$    | U <sup>22</sup> | $U^{33}$    | $U^{12}$     | $U^{13}$     | $U^{23}$     |
|------|-------------|-----------------|-------------|--------------|--------------|--------------|
| C1   | 0.0269 (10) | 0.0383 (12)     | 0.0413 (12) | 0.0072 (9)   | -0.0054 (8)  | -0.0019 (9)  |
| C3   | 0.0358 (11) | 0.0373 (12)     | 0.0398 (12) | 0.0061 (9)   | -0.0028 (9)  | -0.0038 (9)  |
| C10  | 0.0430 (13) | 0.0431 (13)     | 0.0515 (14) | -0.0053 (10) | 0.0050 (10)  | -0.0062 (11) |
| C11  | 0.0496 (14) | 0.0408 (14)     | 0.0692 (17) | -0.0103 (12) | 0.0197 (12)  | -0.0113 (12) |
| C12A | 0.068 (5)   | 0.037 (6)       | 0.092 (5)   | 0.006 (4)    | 0.017 (4)    | -0.010 (4)   |
| C13A | 0.081 (5)   | 0.040 (4)       | 0.109 (9)   | 0.008 (4)    | 0.015 (5)    | -0.023 (4)   |
| C14A | 0.074 (7)   | 0.047 (4)       | 0.114 (9)   | -0.003 (4)   | 0.021 (6)    | -0.036 (5)   |
| C15A | 0.088 (5)   | 0.063 (5)       | 0.083 (5)   | -0.008 (4)   | 0.018 (4)    | -0.040 (4)   |
| C16A | 0.068 (4)   | 0.055 (5)       | 0.071 (4)   | -0.005 (3)   | 0.010 (3)    | -0.029 (3)   |
| C12B | 0.086 (12)  | 0.035 (9)       | 0.108 (14)  | -0.010 (8)   | 0.051 (11)   | -0.003 (9)   |
| C13B | 0.113 (13)  | 0.068 (9)       | 0.112 (15)  | -0.028 (9)   | 0.029 (11)   | -0.039 (11)  |
| C14B | 0.094 (13)  | 0.060 (10)      | 0.095 (15)  | -0.015 (10)  | 0.016 (11)   | -0.033 (10)  |
| C15B | 0.089 (18)  | 0.060 (13)      | 0.090 (12)  | -0.012 (11)  | 0.015 (12)   | -0.014 (11)  |
| C16B | 0.071 (12)  | 0.053 (11)      | 0.103 (11)  | -0.020 (9)   | 0.024 (8)    | -0.012 (9)   |
| C20  | 0.0320 (10) | 0.0471 (13)     | 0.0437 (12) | -0.0021 (10) | 0.0012 (9)   | -0.0005 (10) |
| C21  | 0.0395 (11) | 0.0475 (13)     | 0.0403 (12) | 0.0028 (10)  | 0.0078 (9)   | 0.0014 (10)  |
| C22  | 0.0624 (16) | 0.0777 (19)     | 0.0625 (17) | -0.0047 (14) | -0.0143 (13) | 0.0092 (15)  |
| C23  | 0.070 (2)   | 0.123 (3)       | 0.071 (2)   | 0.007 (2)    | -0.0164 (15) | 0.031 (2)    |
| C24  | 0.082 (2)   | 0.102 (3)       | 0.076 (2)   | 0.029 (2)    | 0.0143 (18)  | 0.040 (2)    |
| C25  | 0.122 (3)   | 0.0551 (19)     | 0.099 (3)   | 0.005 (2)    | 0.015 (2)    | 0.0219 (18)  |
| C26  | 0.083 (2)   | 0.0545 (17)     | 0.073 (2)   | -0.0065 (15) | -0.0062 (16) | 0.0080 (14)  |
|      |             |                 |             |              |              |              |

# supporting information

| N1         | 0.0290 (9) | 0.0431 (11) | 0.0605 (12) | 0.0037 (8) | 0.0001 (8)  | -0.0168 (9) |  |
|------------|------------|-------------|-------------|------------|-------------|-------------|--|
| N2         | 0.0309 (8) | 0.0370 (10) | 0.0355 (9)  | 0.0025 (7) | 0.0006 (7)  | -0.0037 (7) |  |
| 01         | 0.0418 (9) | 0.0742 (12) | 0.0691 (12) | 0.0019 (9) | -0.0104 (8) | -0.0170 (9) |  |
| <b>S</b> 1 | 0.0406 (3) | 0.0480 (4)  | 0.0568 (4)  | 0.0003 (3) | 0.0084 (3)  | 0.0033 (3)  |  |

Geometric parameters (Å, °)

| C1—N2                   | 1.325 (3)   | C12B—H12B      | 0.93        |
|-------------------------|-------------|----------------|-------------|
| C1—N1                   | 1.408 (3)   | C13B—C14B      | 1.361 (15)  |
| C1—S1                   | 1.673 (2)   | C13B—H13B      | 0.93        |
| C3—N2                   | 1.460 (2)   | C14B—C15B      | 1.374 (17)  |
| C3—C3 <sup>i</sup>      | 1.523 (4)   | C14B—H14B      | 0.93        |
| С3—НЗА                  | 0.97        | C15B—C16B      | 1.379 (16)  |
| С3—Н3В                  | 0.97        | C15B—H15B      | 0.93        |
| C10—O1                  | 1.227 (3)   | C16B—H16B      | 0.93        |
| C10—N1                  | 1.370 (3)   | C20—N2         | 1.476 (3)   |
| C10-C11                 | 1.474 (3)   | C20—C21        | 1.504 (3)   |
| C11—C16B                | 1.362 (16)  | C20—H20A       | 0.97        |
| C11—C12A                | 1.386 (7)   | C20—H20B       | 0.97        |
| C11—C12B                | 1.387 (18)  | C21—C26        | 1.373 (3)   |
| C11—C16A                | 1.403 (8)   | C21—C22        | 1.381 (3)   |
| C12A—C13A               | 1.395 (10)  | C22—C23        | 1.382 (4)   |
| C12A—H12A               | 0.93        | С22—Н22        | 0.93        |
| C13A—C14A               | 1.371 (9)   | C23—C24        | 1.343 (5)   |
| C13A—H13A               | 0.93        | С23—Н23        | 0.93        |
| C14A—C15A               | 1.394 (9)   | C24—C25        | 1.356 (4)   |
| C14A—H14A               | 0.93        | C24—H24        | 0.93        |
| C15A—C16A               | 1.399 (9)   | C25—C26        | 1.388 (4)   |
| C15A—H15A               | 0.93        | С25—Н25        | 0.93        |
| C16A—H16A               | 0.93        | C26—H26        | 0.93        |
| C12B—C13B               | 1.388 (17)  | N1—H1N         | 0.861 (10)  |
| N2—C1—N1                | 116.24 (18) | C13B—C14B—C15B | 123 (2)     |
| N2—C1—S1                | 124.30 (16) | C13B—C14B—H14B | 118.5       |
| N1—C1—S1                | 119.40 (16) | C15B—C14B—H14B | 118.5       |
| N2-C3-C3 <sup>i</sup>   | 110.9 (2)   | C14B—C15B—C16B | 118 (2)     |
| N2—C3—H3A               | 109.5       | C14B—C15B—H15B | 120.9       |
| C3 <sup>i</sup> —C3—H3A | 109.5       | C16B—C15B—H15B | 120.9       |
| N2—C3—H3B               | 109.5       | C11—C16B—C15B  | 124 (2)     |
| C3 <sup>i</sup> —C3—H3B | 109.5       | C11—C16B—H16B  | 117.9       |
| НЗА—СЗ—НЗВ              | 108         | C15B—C16B—H16B | 117.9       |
| O1—C10—N1               | 121.0 (2)   | N2-C20-C21     | 111.86 (16) |
| O1—C10—C11              | 122.1 (2)   | N2-C20-H20A    | 109.2       |
| N1-C10-C11              | 116.9 (2)   | C21—C20—H20A   | 109.2       |
| C16B—C11—C12B           | 112 (2)     | N2-C20-H20B    | 109.2       |
| C12A—C11—C16A           | 121.1 (8)   | C21—C20—H20B   | 109.2       |
| C16B—C11—C10            | 132.2 (14)  | H20A-C20-H20B  | 107.9       |
| C12A—C11—C10            | 124.0 (7)   | C26—C21—C22    | 118.0 (2)   |
|                         |             |                | . *         |

| C12B—C11—C10   | 115.6 (18)  | C26—C21—C20  | 121.4 (2)   |
|----------------|-------------|--------------|-------------|
| C16A—C11—C10   | 114.7 (6)   | C22—C21—C20  | 120.5 (2)   |
| C11—C12A—C13A  | 118.2 (10)  | C21—C22—C23  | 120.2 (3)   |
| C11—C12A—H12A  | 120.9       | C21—C22—H22  | 119.9       |
| C13A—C12A—H12A | 120.9       | C23—C22—H22  | 119.9       |
| C14A—C13A—C12A | 121.7 (8)   | C24—C23—C22  | 121.2 (3)   |
| C14A—C13A—H13A | 119.1       | С24—С23—Н23  | 119.4       |
| C12A—C13A—H13A | 119.1       | С22—С23—Н23  | 119.4       |
| C13A—C14A—C15A | 120.3 (7)   | C23—C24—C25  | 119.5 (3)   |
| C13A—C14A—H14A | 119.9       | C23—C24—H24  | 120.2       |
| C15A—C14A—H14A | 119.9       | C25—C24—H24  | 120.2       |
| C14A—C15A—C16A | 119.3 (8)   | C24—C25—C26  | 120.4 (3)   |
| C14A—C15A—H15A | 120.3       | С24—С25—Н25  | 119.8       |
| C16A—C15A—H15A | 120.3       | С26—С25—Н25  | 119.8       |
| C15A—C16A—C11  | 119.4 (8)   | C21—C26—C25  | 120.5 (3)   |
| C15A—C16A—H16A | 120.3       | C21—C26—H26  | 119.7       |
| C11—C16A—H16A  | 120.3       | C25—C26—H26  | 119.7       |
| C11—C12B—C13B  | 129 (4)     | C10—N1—C1    | 122.61 (17) |
| C11—C12B—H12B  | 115.6       | C10—N1—H1N   | 121.7 (16)  |
| C13B—C12B—H12B | 115.6       | C1—N1—H1N    | 115.6 (16)  |
| C14B—C13B—C12B | 113 (3)     | C1—N2—C3     | 120.25 (17) |
| C14B—C13B—H13B | 123.5       | C1—N2—C20    | 125.18 (17) |
| C12B—C13B—H13B | 123.5       | C3—N2—C20    | 114.56 (16) |
|                |             |              |             |
| O1—C10—C1—S1   | -62.55 (17) | S1-C1-N1-C10 | -49.1 (3)   |
| O1-C10-N1-C1   | -24.3 (3)   |              |             |
|                |             |              |             |

Symmetry code: (i) -x+2, -y+1, -z+1.

Hydrogen-bond geometry (Å, °)

| D—H···A                             | D—H      | H···A    | D···A     | D—H···A |
|-------------------------------------|----------|----------|-----------|---------|
| N1—H1 <i>N</i> …O1 <sup>ii</sup>    | 0.86(1)  | 2.30(1)  | 3.073 (2) | 150 (2) |
| N1—H1 <i>N</i> ····S1 <sup>ii</sup> | 0.86 (1) | 2.98 (2) | 3.647 (2) | 136 (2) |

Symmetry code: (ii) *x*–1, *y*, *z*.