



Acta Crystallographica Section E

Structure Reports

Online

ISSN 1600-5368

4-Bromo-N-(4-bromophenyl)aniline

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Received 14 February 2011; accepted 24 February 2011

Key indicators: single-crystal X-ray study; T = 125 K; mean $\sigma(C-C) = 0.003$ Å; R factor = 0.026; wR factor = 0.064; data-to-parameter ratio = 24.3.

In the title compound, $C_{12}H_9Br_2N$, the dihedral angle between the benzene rings is 47.32 (5)°, whereas the pitch angles, or the angles between the mean plane of each aryl group 'propeller blade' and the plane defined by the aryl bridging C-N-C angle, are 18.1 (2) and 31.7 (2)°. No intermolecular N-H hydrogen bonding is present in the crystal; however, there is a short intermolecular $Br \cdots Br$ contact of 3.568 (1) Å.

Related literature

The title compound is an amine analogue of brominated diphenyl ether flame retardant materials commonly used in household items. For information on environmental and health concerns related to brominated flame retardants, see: de Wit (2002); Lunder *et al.* (2010). For the synthesis of the title compound, see: Crounse & Raiford (1945); Galatis & Megaloikonomos (1934); He *et al.* (2008). For related structures, see: Eriksson *et al.* (2004); Plieth & Ruban (1961); Li *et al.* (2010). For the van der Waals radius of Br and intermolecular Br···Br contacts, see: Bondi (1964); Medlycott *et al.* (2007). For a description of the pitch angle, see: Lim & Tanski (2007).

Experimental

Crystal data

 $C_{12}H_9Br_2N$

 $M_r = 327.02$

Monoclinic, $P2_1/c$ Z = 4 Mo $K\alpha$ radiation b = 13.032 (3) Å $\mu = 7.30 \text{ mm}^{-1}$ c = 14.228 (3) Å T = 125 K $\beta = 96.967$ (3)° V = 1104.2 (4) Å³

Data collection

Bruker APEXII CCD diffractometer 3373 independent reflections 3786 reflections with $I > 2\sigma(I)$ $R_{\rm int} = 0.218, \ T_{\rm max} = 0.370$

Refinement

 $\begin{array}{ll} R[F^2>2\sigma(F^2)]=0.026 & \text{H atoms treated by a mixture of} \\ wR(F^2)=0.064 & \text{independent and constrained} \\ S=1.02 & \text{refinement} \\ 3373 \text{ reflections} & \Delta\rho_{\max}=0.94 \text{ e Å}^{-3} \\ 139 \text{ parameters} & \Delta\rho_{\min}=-0.45 \text{ e Å}^{-3} \end{array}$

Data collection: *APEX2* (Bruker, 2007); cell refinement: *SAINT* (Bruker, 2007); data reduction: *SAINT*; program(s) used to solve structure: *SHELXS97* (Sheldrick, 2008); program(s) used to refine structure: *SHELXL97* (Sheldrick, 2008); molecular graphics: *SHELXTL* (Sheldrick, 2008); software used to prepare material for publication: *SHELXTL*.

This work was supported by Vassar College. X-ray facilities were provided by the US National Science Foundation (grant No. 0521237 to JMT).

Supplementary data and figures for this paper are available from the IUCr electronic archives (Reference: SI2339).

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Acta Cryst. (2011). E67, o755 [doi:10.1107/S160053681100715X]

4-Bromo-N-(4-bromophenyl)aniline

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S1. Comment

The title compound, 4-bromo-*N*-(4-bromophenyl)aniline, C₁₂H₉Br₂N (I), was first synthesized by Galatis & Megaloikonomos (1934) *via* the direct bromination of diphenylamine, and the structure was corroborated by Crounse & Raiford (1945) in their study of the hydrolysis of the benzoyl derivative. More recently, halogenated diphenylamines have been prepared by copper catalyzed coupling reactions (He *et al.*, 2008). The crystal structure of the chloride analogue is known (Plieth & Ruban, 1961), and an analogous structure with an oxygen bridge has also been reported (Eriksson *et al.*, 2004). The title compound is an amine analogues of a class of brominated diphenyl ether materials (de Wit, 2002). Polybrominated diphenyl ethers are commonly used as flame retardants (Eriksson *et al.*, 2004) in consumer products and electronics and have been found in humans (Lunder *et al.*, 2010).

Compound (I) is a dibrominated diphenyl amine derivative with a "propeller blade" disposition of the benzene rings about the bridging nitrogen atom. The structure reveals that there is no intermolecular hydrogen bonding, although there are significant intermolecular Br···Br contacts (Medlycott *et al.*, 2007) at a distance of 3.568 (1) Å, which is shorter than the sum of the van der Waals radius of bromine, 1.85Å (Bondi, 1964), at 3.7 Å. The aryl-bridging C4—N—C7 angle in (I) is 128.5 (2)°, somewhat smaller than the C—N—C bond angle of 133.8° found in the isomorphous dichloro analog (Plieth & Ruban, 1961), but similar to the C—N—C bond angle of 128.1° in another similar structure, *N*-4-(bromophenyl)-4-nitroaniline, which contains one bromo and one nitro group (Li *et al.*, 2010).

The dihedral angle in (I) is found to be 47.32 (5)°, whereas the pitch angles are 18.1 (2)° and 31.7 (2)°. The pitch angles are the angles between the mean plane of each aryl group "propeller blade" and the plane defined by the aryl bridging C4—N—C7 angle. The pitch angles are metrical parameters that describe the dispostion of the benzene rings about the bridging atom with greater detail than the dihedral angle; structures with equivalent dihedral angles may exhibit dramatically different orientations of the benzene rings about the bridging group (Lim & Tanski, 2007). In the isomorphous dichloro analog to the title compound, the dihedral angle is found to be significantly larger, 56.5°, as are the pitch angles of 22.1° and 39.1°. In another similar bromo compound, *N*-4-(bromophenyl)-4-nitroaniline, where the dihedral angle of 44.8° is more similar to that of the title compound, the pitch angles are found to be 12.6° and 35.1°.

S2. Experimental

Crystalline 4-bromo-N-(4-bromophenyl)aniline (I) was purchase from Aldrich Chemical Company, USA.

S3. Refinement

All non-hydrogen atoms were refined anisotropically. The hydrogen atoms on carbon were included in calculated positions and were refined using a riding model at C–H = 0.95Å and $U_{iso}(H) = 1.2 \times U_{eq}(C)$ of the aryl C-atoms. T hydrogen atom on nitrogen was refined semifreely with the help of a distance restraint, d(N-H) = 0.835 (16) Å and $U_{iso}(H) = 1.2 \times U_{eq}(N)$. The extinction parameter (EXTI) refined to zero and was removed from the refinement.

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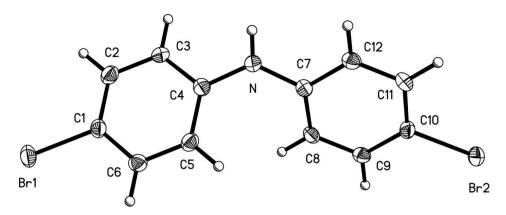


Figure 1

A view of compound (I), with displacement ellipsoids shown at the 50% probability level.

4-Bromo-N-(4-bromophenyl)aniline

Crystal data

C12H9Br2N $M_r = 327.02$ Monoclinic, $P2_1/c$ Hall symbol: -P 2vbc a = 5.9993 (12) Å b = 13.032 (3) Å c = 14.228 (3) Å $\beta = 96.967 (3)^{\circ}$ $V = 1104.2 (4) \text{ Å}^3$

Z=4

Data collection

Bruker APEXII CCD diffractometer

Radiation source: fine-focus sealed tube

Graphite monochromator

 φ and ω scans

Absorption correction: multi-scan (SADABS; Bruker 2007) $T_{\min} = 0.218, T_{\max} = 0.370$

Refinement

Refinement on F^2 Least-squares matrix: full $R[F^2 > 2\sigma(F^2)] = 0.026$ $wR(F^2) = 0.064$ S = 1.02

3373 reflections 139 parameters 1 restraint

Primary atom site location: structure-invariant

direct methods

F(000) = 632 $D_{\rm x} = 1.967 \; {\rm Mg \; m^{-3}}$

Mo $K\alpha$ radiation, $\lambda = 0.71073 \text{ Å}$ Cell parameters from 8371 reflections

 $\theta = 2.9 - 30.4^{\circ}$

 $\mu = 7.30 \text{ mm}^{-1}$

T = 125 K

Plate, colourless

 $0.30 \times 0.30 \times 0.17 \text{ mm}$

17275 measured reflections 3373 independent reflections 2786 reflections with $I > 2\sigma(I)$

 $R_{\rm int} = 0.038$

 $\theta_{\text{max}} = 30.6^{\circ}, \, \theta_{\text{min}} = 2.1^{\circ}$

 $h = -8 \longrightarrow 8$

 $k = -18 \rightarrow 18$

 $l = -20 \rightarrow 20$

Secondary atom site location: difference Fourier

Hydrogen site location: inferred from neighbouring sites

H atoms treated by a mixture of independent and constrained refinement

 $w = 1/[\sigma^2(F_0^2) + (0.0365P)^2]$ where $P = (F_0^2 + 2F_c^2)/3$

 $(\Delta/\sigma)_{\text{max}} = 0.001$

 $\Delta \rho_{\rm max} = 0.94 \text{ e Å}^{-3}$

 $\Delta \rho_{\min} = -0.45 \text{ e Å}^{-3}$

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Special details

Experimental. A suitable crystal was mounted in a nylon loop with Paratone-N cryoprotectant oil and data was collected on a Bruker *APEX* 2 CCD platform diffractometer. The structure was solved using direct methods and standard difference map techniques, and was refined by full-matrix least-squares procedures on F² with *SHELXTL* Version 6.14 (Sheldrick, 2008).

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 F-factors based on ALL data will be even larger.

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

	x	y	Z	$U_{ m iso}$ */ $U_{ m eq}$
Br1	0.35642 (4)	0.125752 (16)	0.143618 (15)	0.02649 (7)
Br2	-0.19656(3)	0.946875 (15)	0.134200 (15)	0.02372 (7)
N	0.4407 (3)	0.58823 (13)	0.10486 (13)	0.0224 (4)
H1	0.566 (3)	0.6043 (18)	0.0893 (16)	0.027*
C1	0.3790(3)	0.27103 (16)	0.13498 (13)	0.0200 (4)
C2	0.5839(3)	0.31866 (15)	0.16241 (14)	0.0229 (4)
H2A	0.7114	0.2793	0.1865	0.027*
C3	0.5998 (3)	0.42427 (15)	0.15413 (14)	0.0215 (4)
H3A	0.7391	0.4573	0.1733	0.026*
C4	0.4131 (3)	0.48316 (15)	0.11792 (13)	0.0186 (4)
C5	0.2092(3)	0.43296 (15)	0.09209 (14)	0.0204 (4)
H5A	0.0806	0.4719	0.0685	0.025*
C6	0.1911 (3)	0.32763 (15)	0.10030 (14)	0.0209 (4)
H6A	0.0514	0.2944	0.0824	0.025*
C7	0.2844 (3)	0.66694 (15)	0.10980 (14)	0.0188 (4)
C8	0.0933 (3)	0.65683 (15)	0.15638 (14)	0.0192 (4)
H8A	0.0621	0.5931	0.1845	0.023*
C9	-0.0513(3)	0.73940 (15)	0.16173 (13)	0.0194 (4)
H9A	-0.1822	0.7318	0.1925	0.023*
C10	-0.0044(3)	0.83266 (14)	0.12218 (14)	0.0191 (4)
C11	0.1859 (3)	0.84517 (15)	0.07631 (14)	0.0208 (4)
H11A	0.2179	0.9096	0.0497	0.025*
C12	0.3281 (3)	0.76224 (15)	0.06992 (13)	0.0202 (4)
H12A	0.4572	0.7701	0.0380	0.024*

Atomic displacement parameters (\mathring{A}^2)

	U^{11}	U^{22}	U^{33}	U^{12}	U^{13}	U^{23}
Br1	0.03250 (12)	0.01599 (11)	0.03174 (12)	0.00176 (8)	0.00695 (9)	-0.00159 (8)
Br2	0.02408 (11)	0.01647 (10)	0.03091 (12)	0.00214 (7)	0.00456 (8)	-0.00193 (7)
N	0.0179 (8)	0.0174 (8)	0.0330 (10)	0.0004(7)	0.0076 (7)	0.0013 (7)
C1	0.0219 (9)	0.0166 (9)	0.0217 (9)	0.0016 (7)	0.0039 (7)	-0.0020(7)

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C2	0.0197 (9)	0.0232 (10)	0.0253 (10)	0.0046 (8)	0.0008 (8)	0.0007 (8)
C3	0.0144 (9)	0.0215 (10)	0.0283 (10)	-0.0001(7)	0.0012 (7)	-0.0016 (8)
C4	0.0203 (9)	0.0182 (9)	0.0180 (9)	-0.0002(7)	0.0051(7)	-0.0009(7)
C5	0.0184 (9)	0.0210 (10)	0.0212 (9)	0.0031 (7)	-0.0006(7)	-0.0014 (7)
C6	0.0189 (9)	0.0225 (10)	0.0211 (9)	-0.0008(7)	0.0020(7)	-0.0033 (7)
C7	0.0192 (9)	0.0167 (9)	0.0205 (9)	-0.0008(7)	0.0019(7)	0.0001 (7)
C8	0.0204 (9)	0.0155 (9)	0.0219 (9)	-0.0030(7)	0.0038 (7)	0.0002 (7)
C9	0.0176 (9)	0.0207 (10)	0.0206 (9)	-0.0031(7)	0.0049 (7)	-0.0008(7)
C10	0.0192 (9)	0.0150 (9)	0.0227 (9)	0.0017(7)	0.0005 (7)	-0.0019 (7)
C11	0.0239 (10)	0.0156 (9)	0.0232 (10)	-0.0042(7)	0.0038 (8)	0.0009 (7)
C12	0.0193 (9)	0.0210 (9)	0.0210 (9)	-0.0027(7)	0.0049 (7)	-0.0007(7)

Geometric parameters (Å, °)

Br1—C1	1.903 (2)	С5—Н5А	0.9500
Br2—C10	1.9031 (19)	C6—H6A	0.9500
N—C4	1.394 (3)	C7—C8	1.399 (3)
N—C7	1.397 (3)	C7—C12	1.403 (3)
N—H1	0.835 (16)	C8—C9	1.390 (3)
C1—C6	1.387 (3)	C8—H8A	0.9500
C1—C2	1.390 (3)	C9—C10	1.383 (3)
C2—C3	1.386 (3)	С9—Н9А	0.9500
C2—H2A	0.9500	C10—C11	1.391 (3)
C3—C4	1.403 (3)	C11—C12	1.387 (3)
С3—Н3А	0.9500	C11—H11A	0.9500
C4—C5	1.397 (3)	C12—H12A	0.9500
C5—C6	1.383 (3)		
C4—N—C7	128.53 (17)	C1—C6—H6A	120.4
C4—N—H1	114.0 (17)	N—C7—C8	123.26 (17)
C7—N—H1	117.4 (17)	N—C7—C12	118.05 (17)
C6—C1—C2	121.03 (19)	C8—C7—C12	118.62 (18)
C6—C1—Br1	119.41 (15)	C9—C8—C7	120.40 (18)
C2—C1—Br1	119.56 (15)	C9—C8—H8A	119.8
C3—C2—C1	119.14 (18)	C7—C8—H8A	119.8
C3—C2—H2A	120.4	C10—C9—C8	119.93 (18)
C1—C2—H2A	120.4	C10—C9—H9A	120.0
C2—C3—C4	120.97 (18)	C8—C9—H9A	120.0
C2—C3—H3A	119.5	C9—C10—C11	120.87 (18)
C4—C3—H3A	119.5	C9—C10—Br2	119.70 (15)
N—C4—C5	122.64 (18)	C11—C10—Br2	119.40 (14)
N—C4—C3	118.93 (17)	C12—C11—C10	119.05 (18)
C5—C4—C3	118.36 (18)	C12—C11—H11A	120.5
C6—C5—C4	121.20 (18)	C10—C11—H11A	120.5
C6—C5—H5A	119.4	C11—C12—C7	121.13 (18)
C4—C5—H5A	119.4	C11—C12—H12A	119.4
C5—C6—C1	119.28 (19)	C7—C12—H12A	119.4
C5—C6—H6A	120.4		

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-0.5(3)	C4—N—C7—C8	20.0(3)
178.75 (15)	C4—N—C7—C12	-163.25 (19)
-0.5(3)	N—C7—C8—C9	177.59 (18)
33.3 (3)	C12—C7—C8—C9	0.8 (3)
-149.9(2)	C7—C8—C9—C10	-1.1(3)
-175.67 (18)	C8—C9—C10—C11	0.4(3)
1.3 (3)	C8—C9—C10—Br2	-177.49 (14)
175.75 (18)	C9—C10—C11—C12	0.5(3)
-1.1(3)	Br2—C10—C11—C12	178.40 (15)
0.1 (3)	C10—C11—C12—C7	-0.8(3)
0.7 (3)	N—C7—C12—C11	-176.84 (18)
-178.55 (14)	C8—C7—C12—C11	0.1 (3)
	178.75 (15) -0.5 (3) 33.3 (3) -149.9 (2) -175.67 (18) 1.3 (3) 175.75 (18) -1.1 (3) 0.1 (3) 0.7 (3)	178.75 (15)

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