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Boron monoxide is a one-dimensional polymer

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It was recently reported that boron monoxide (BO) is formed through the cross-linking of B₄O₂ structural building units. Multiple theoretical phases agree with this description. Using pycnometry, multidimensional ¹⁷O NMR spectroscopy, and plane-wave DFT calculations we determined the likely polymorph to be a onedimensional polymer initially proposed in 1955.

Among the binary oxides, boron monoxide (BO) is perhaps unique in having evaded attempts to determine its structure for nearly a century. The material was initially reported in 1940, 1,2 however, the first synthesis for the preparation of a singlephase BO material was described in 1955 by Wartik and Apple.³ Their synthesis employed the condensation of tetrahydroxydiboron (B₂(OH)₄, Fig. 1) at high temperature and while it was soon discovered that the boron-boron bonds were preserved in the resulting material, 4,5 the long-range structure was never determined. Several potential model structures were reported, 6-8 most exhaustively by Claeyssens et al.,9 however, none were supported by experimental evidence.

Recently, some of us applied multidimensional ¹¹B NMR spectroscopy to revisit this structural conundrum. 10 Key observations included (1) the detection of a single crystallographically unique boron site; (2) observation of symmetry-amplified ¹¹B-¹¹B J couplings; ¹¹⁻¹³ (3) observation of the collinearity of closest B-B bonds from 11B-11B-11B triple-quantum correlations; and (4) observation of diffraction signals that suggested the formation of a layered structure. The work strongly suggested that the main structural building unit was a B₄O₂ ring with local D_{2h} symmetry. ¹⁰ This agrees with the isolation of a B₄O₂(OH)₄ intermediate by Carmalt et al. 14 Owing to the observed diffraction patterns we proposed that the material

Fig. 1 The synthesis of BO involves the condensation of B₂(OH)₄ to form structural building units composed of a six-membered B₄O₂ rings that then interlink into a polymeric material. The structural building units can organize into either one-dimensional (A), two-dimensional (B) and (C), or three-dimensional (D) and (E) structures. Three-dimensional structures (D) and (E) feature twisting of the B₂O₄ moieties, which is inconsistent with the observation of symmetry-amplified J splittings. Darker shades indicate moieties in the foreground

was two-dimensional, which could have important practical implications; 15-21 however, a great deal of uncertainty on the structure remained. B₄O₂ rings may cross-link in a number of patterns, as evident from the structures compiled by Claeyssens and listed in Fig. 1.9 We thus sought to apply 17O NMR spectroscopy to further narrow the possible structures for BO. While most materials are indistinguishable from ¹¹B NMR (with the exception of structure D and E, which do not have local D_{2h} symmetry), structure **A** features a single unique ¹⁷O site while the others feature intraring and interring oxygens that may be distinguishable. All five models further differ in their B-B-O-B dihedral angles that could be probed using ¹⁷O{¹¹B} heteronuclear correlation.

^{*}B₂O₄ distorsions inconsistent with J splitting

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To this aim, we enriched B₂(OH)₄ with ¹⁷O to prepare an ¹⁷O-enriched BO material. Enrichment of tetrahydroxydiboron by direct ¹⁷O exchange with water proved ineffective due to its poor solubility and slow decomposition to form boric acid, a non-isolable impurity. We therefore instead produced tetramethoxydiboron as previously described, 22 which that then hydrolyzed with an excess of ¹⁷O-enriched water (39.9%) to vield pure B₂(¹⁷OH)₄. From liquid-state ¹⁷O NMR, we estimate the enrichment level to be $\sim 3.9\%$ (Fig. S5). Synthesis of B¹⁷O was then carried out in the usual fashion by the self-condensation of $B_2(^{17}OH)_4$ at 200 °C, see SI for further details.

We acquired 11B and 17O multiple-quantum (MQ) magicangle spinning (MAS) NMR spectra of the resulting material (Fig. 2). 23-25 As expected, 10 we observed a single 11B resonance with an isotropic (δ_{iso}) chemical shift of 35.1 ppm, a quadrupolar coupling constant (C_0) of 3.5 MHz, and a quadrupolar asymmetry parameter (η) of 0.5. The ¹⁷O MQMAS spectrum was equally simple, being described with a single site (δ_{iso} = 178 ppm; $C_{\rm O}$ = 3.8 MHz; η = 0.7, see Table 1). These parameters were obtained by fitting the MAS lineshape and the MQMAS shift simultaneously using dmfit.²⁶

Immediately, these results strongly suggest the formation of structure A (see Fig. 1), which uniquely features only one type of chemically distinct oxygen site. We then performed plane-wave density functional theory (DFT) calculations to predict the ¹¹B and ¹⁷O chemical shifts and electric field gradient (EFG) tensor^{27,28} so that they may be compared with experiment. We found that the ¹⁷O chemical shifts of sites within ring structures were highly sensitive to the method used, and as such a monomer correction29 was applied using the resolution of identity second-order Møller-Plesset perturbation theory method (RI-MP2) for all calculated ¹⁷O chemical shifts. The results are listed in Table 1. There is a clear distinction between the intraring (δ_{iso} = 200–225 ppm) and interring oxygen species ($\delta_{\rm iso}$ = 150-165 ppm) suggesting that structure **B-F** are inconsistent with the ¹⁷O NMR observations.

The differences between the various model are perhaps most clearly depicted by comparing experimental and computed ¹⁷O{¹¹B} dipolar heteronuclear multiple-quantum correlation

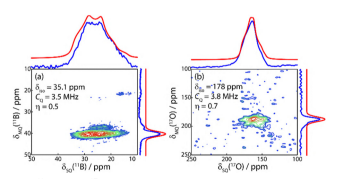


Fig. 2 11 B (a) and 17 O (b) 2D MQMAS NMR spectra. There is one welldefined site present for both ¹¹B and ¹⁷O, suggesting a structure containing only a single crystallographically unique site for each element. Sum projections (blue) are overlaid with a fit to a single site with the listed chemical shift and EFG tensor parameters (red)

Table 1 Experimentally-determined and computed BO NMR parameters and densities

	$ ho/{ m g~cm}^{-3}$	Site	$\delta_{\rm iso}$ /ppm	$ C_{\rm Q} /{ m MHz}$	η
Expt.	2.08 ± 0.13	В	35 ± 1	3.5 ± 0.1	0.5 ± 0.2
•		O	178 ± 2	3.8 ± 0.1	0.7 ± 0.1
\mathbf{A}^{a}	2.05	В	33.8	3.90	0.32
		O	203.9	4.73	0.52
В	1.53	В	28.2	3.86	0.44
		O_{intra}	223.1	5.16	0.35
		O_{inter}	149.8	6.87	0.51
C	1.44	В	25.5	3.65	0.50
		В	31.0	3.95	0.38
		O_{intra}	217.0	4.84	0.50
		O_{intra}	219.4	4.48	0.66
		O_{inter}	161.3	5.03	0.81
D	0.84	В	31.7	3.97	0.38
		В	25.5	5.66	0.95
		O_{intra}	215.9	4.48	0.70
		O_{intra}	205.4	4.31	0.40
		O_{inter}	163.2	4.79	0.30
		O_{inter}	162.2	4.84	0.36
E	1.35	В	30.9	4.28	0.64
		В	27.7	6.18	0.94
		O_{intra}	213.6	4.60	0.63
		O_{intra}	218.8	4.67	0.26
		O_{inter}	153.6	4.81	0.95

^a Calculation performed on the *Immm* polymorph.⁹

(D-HMQC)³⁰ spectra (Fig. 3) that depend on the ¹¹B and ¹⁷O EFG tensors (including their relative orientations) in addition to the chemical shifts. We acquired such a spectrum using rotational-echo double-resonance (REDOR) recoupling,31 applied at the 11B frequency, and 17O detection. 32,33 Owing to the strong 11B-11B homonuclear dipolar interactions and the low ¹⁷O concentration, ¹¹B-detection was not feasible. DFTpredicted ¹⁷O{¹¹B} correlation spectra were calculated using SIMPSON^{34,35} with the parameters listed in Table 1, Euler angles calculated using MagresView ver. 1.6.2 (Table S2),36 and assuming a 100% efficient magnetization transfer. Clearly only structure A yields a correlation spectrum in close agreement with that measured experimentally, with the others having clearly defined intraring and interring correlations. Three-dimensional structures D and E feature ring distortions that disagree with prior I coupling measurements. 10,11 These distortions further dramatically increase the magnitude of the ¹¹B quadrupolar interactions beyond that observed experimentally. Comparisons between the 1D Hahn echo NMR spectra and the five models are also shown in Fig. S2.

To further confirm that structure **A** is the correct polymorph for the BO materials produced by the condensation of B₂(OH)₄, we performed pycnometry measurements to determine the density of the material. As described by Claevssens, and as listed in Table 1, the polymorphs differ greatly in their densities, with structure **A** being the densest at a predicted 2.05 g cm^{-3} . We measured a density of 2.08 \pm 0.13, which is within error of the predicted value for A, and far denser than structures B-E.

It thus seems that structure A, a one-dimensional polymer, is indeed the correct model for BO, however, this does not explain the peculiar X-ray reflections that pointed to a twodimensional material. Claeyssens predicted four different



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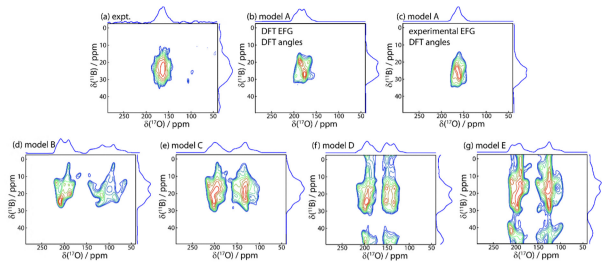


Fig. 3 Experimental (a) and calculated (b)-(q) ¹⁷O(¹¹B) D-HMQC 2D correlation spectra. Simulations were performed using the periodic DFT-calculated chemical shifts and EFG tensor parameters for the structures indicated on the Figure. For the simulation shown in (c), the Euler angles from the DFT calculations were used together with the experimentally-determined EFG tensor parameters and chemical shifts.

packing arrangements for the polymers in A,9 which lead to very different diffraction patterns (Fig. 4), none of which agree with the experimental result. If we randomize the stacking arrangement in a 1 \times 1 \times 10 P1 supercell, 10 however, we do reproduce a similar powder pattern as that predicted for structure B. There are still differences with the experiment, however, and a broader study of the stacking faults may be required to fully understand the diffraction pattern.

To conclude, density measurements and ¹⁷O{¹¹B} solid-state NMR experiments were used to narrow the potential structural candidates for boron monoxide (BO), which was earlier shown to be made of interconnected B₄O₂ rings. We discovered that the material was, in fact, not a two-dimensional nanomaterial but instead was composed of one-dimensional polymers composed of fused B₄O₂ rings. Interestingly, this exact model

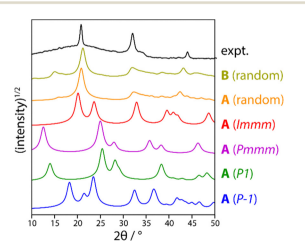


Fig. 4 Comparison of the experimentally-measured PXRD pattern to those predicted using different polymorphs of model ${\bf A}$, as indicated on the Figure, including turbostratic models of A and B. None of the predicted patterns agree with the experimental measurement, suggesting that some degree of random stacking is common.

was initially suggested by Wartick and Apple in 1955 when the material was first prepared.3 We were nevertheless unable to narrow the possibilities for a space group, with the material seeming to lack long-range order. Learning about the longrange order of the material will require methods that are sensitive to such length scales, such as scanning tunneling microscopy.37

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Conflicts of interest

There are no conflicts to declare.

Data availability

Raw data for this study are available at https://doi.org/10.5281/ zenodo.16884299.

Supplementary information (SI): synthesis details, solutionphase NMR data, differential scanning calorimetry, 11B J-resolved NMR data, thermogravimetric analysis, infrared spectroscopy, and further calculation details. See DOI: https://doi.org/ 10.1039/d5cc04723d.

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