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# Ligand solid-solution tuning of magnetic and mechanical properties of the van der Waals metal-organic magnet $NiCl_2(btd)_{1-x}(bod)_x$ †

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Van der Waals (vdW) magnets offer unique opportunities for exploring magnetism in the 2D limit. Metal-organic magnets (MOM) are of particular interest as the functionalisation of organic ligands can control their physical properties. Here, we demonstrate tuning of mechanical and magnetic function of a noncollinear vdW ferromagnet, NiCl<sub>2</sub>(btd) (btd = 2,1,3-benzothiadiazole), through creating solid-solutions with the oxygen-substituted analogue ligand 2,1,3benzoxadiazole (bod). We synthesise  $NiCl_2(btd)_{1-x}(bod)_x$  up to x =0.33, above which we find mixtures primarily composed of 1D NiCl<sub>2</sub>(bod)<sub>2</sub>. Magnetometry reveals bod incorporation reduces the coercivity significantly (up to 60%), without altering the ordering temperatures. High pressure synchrotron diffraction measurements up to 0.4 GPa demonstrate that the stiffest axis is the b axis, through the Ni-N-(O/S)-N-Ni bonds, and the softest is the interlayer direction. Doping with bod fine-tunes this compressibility, softening the layers, but stiffening the interlayer axis. This demonstrates that substitution of organic ligands in vdW MOMs can be used to realise targeted magnetic and mechanical properties.

The modularity of metal–organic materials means that compounds with identical structural topologies but different ligands can be readily synthesised (they are 'isoreticular'). This in turn enables the synthesis of diverse ligand solid-solutions, <sup>2,3</sup> which allows control of chemical function, *e.g.* methane separation and catalytic activity. The physical properties of metal–organic frameworks (MOFs), *e.g.* mechanical, magnetic, or electronic function, can

Ligand solid-solution control over mechanical and magnetic function in vdW magnets is of special interest because pressureand strain-control over magnetism can be readily achieved in devices. This is particularly true for noncollinear magnets, where continuous evolution of magnetic order and properties is possible. We have recently reported a family of new layered MOMs with noncollinear magnetic structures, including the canted ferromagnet NiCl<sub>2</sub>(btd). This material consists of NiCl<sub>2</sub> chains coordinated by the nitrogens of the nonlinear btd ligand to form corrugated sheets [Fig. 1]. The easy-axis ferromagnetic chains in combination with the tilting of chains induced by the ligand geometry leads to noncollinear canted ferromagnetism with significant coercive field,  $\mu_0H_c=1.0(1)$  T. The modularity of this

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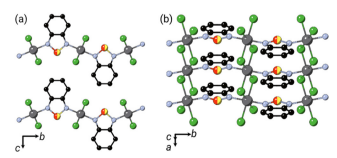


Fig. 1 Structure of  $NiCl_2(btd)_{1-x}(bod)_x$ . (a) Viewed along the [100] direction (b) viewed along the [101] direction. C = black; Ni = grey; Cl = green; N = blue; O/S = red/yellow and H atoms omitted for clarity.

equally be controlled through ligand solution. There remains a great deal to learn about the physical properties of mixed-ligand MOFs, especially the possibility of creating function beyond the linear combination of stoichiometric end-members. Ligand solid-solutions in zeolitic imidazolate frameworks (ZIFs) subtly modulate the magnetic ordering temperatures of **sod** topology ZIFs and control the pressure-induced pore closing ZIF-4 analogues; and solid-solutions of terminal halide ligands in  $Cr(pyz)_2Br_xI_{2-x}$  produce temperature-induced valence tautomeric transitions not present in the stoichiometric phases.

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system, together with the promise of its magnetic function, prompted us to investigate whether we could use substitution of btd ligand for bod to not only alter the structure, but also tune the magnetic and mechanical properties of MOMs.

We showed that NiCl<sub>2</sub>(btd) can be made phase pure and crystalline through the direct reaction of NiCl<sub>2</sub>·6H<sub>2</sub>O and btd, <sup>16</sup> and thus we first explored this approach to create the solidsolutions NiCl<sub>2</sub>(btd)<sub>1-x</sub>(bod)<sub>x</sub>, with target bod fraction,  $x_t = 0$ , 0.25, 0.5, 0.75 and 1.0 [Section S1, ESI†].

Analysis of the powder X-ray diffraction (PXRD) data confirmed that we were able to produce the desired phase up to  $x_t \le 0.75$  [Fig. 2 and Fig. S2, S3, ESI†]. We found that the pure bod phase did not form. Consequently, we synthesised a series through the reaction of ethanolic solutions of nickel chloride and ligands over the same target range of  $x_t$ , analogous to CoCl<sub>2</sub>(btd).<sup>17</sup> We found by analysis of PXRD data this again produced powders isostructural to NiCl<sub>2</sub>(btd) up to  $x_t \le 0.75$ , but at  $x_t = 1$  we obtained a phase mixture with an unknown primary phase. Pawley refinement showed that the samples synthesised through direct reaction contained very small quantities of starting material, but that the solution-synthesised samples had broader diffraction peaks, likely due to small particle sizes [Fig. S2 and Section S2.1, ESI†].

The phase mixture formed during solution synthesis with  $x_t = 1$  included a number of small single crystals [Sections S2.3 and S2.4, ESI†]. We found using single crystal X-ray diffraction these to be a new 1D coordination polymer NiCl2(bod)2, containing trans-NiCl<sub>4</sub>N<sub>2</sub> octahedra connected into NiCl<sub>2</sub> chains with terminal bod ligands [Section S2.3 and Fig. S15, S16, Table S5, ESI†], and a small number of crystals of NiCl<sub>2</sub>·2H<sub>2</sub>O.<sup>18</sup> Re-analysis of our PXRD data in the light of this showed it was primarily NiCl<sub>2</sub>(bod)<sub>2</sub> and a small quantity of nickel chloride hydrates. Further examination using single crystal electron diffraction (SCED) of the remainder of the reaction mixture revealed that the sample contained a number of different phases with unit cells closely related to NiCl<sub>2</sub>(btd), though with slightly different symmetries [Section S2.4 and Fig. S17-S20, Table S6, ESI†].16 Comparison of refinements with only bod, only btd, and mixed ligands showed that these nanocrystals were monoclinic twinned NiCl<sub>2</sub>(bod) and an orthorhombic

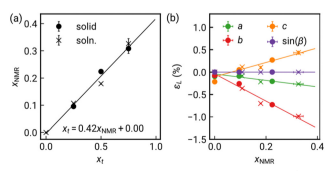


Fig. 2 (a) Determination of x through integration of solution  $^{1}H$  NMR spectra. Linear fit shown. (b) Variation in lattice parameter strain  $(\varepsilon_L)$  with linear fit. Data point for  $x_{NMR} = 0.31$  solid-state excluded due to presence of NiCl<sub>2</sub>·6H<sub>2</sub>O impurity. Circles indicate solid-state samples, crosses solution samples.

polymorph of NiCl<sub>2</sub>(btd), although we cannot exclude that this orthorhombic phase includes a low proportion of bod (<5%). We note these phases are not seen in the bulk PXRD and hence, we ascribe the formation of a small number of nanocrystals of NiCl<sub>2</sub>(btd) to the presence of adventitious btd, likely facilitated by its high vapour pressure [Fig. S4, ESI†]. This highlights the capability of SCED to find and solve the structures of even minor crystalline phases.

We determined the bod content by solution  $^{1}$ H NMR,  $x_{\text{NMR}}$ , by dissolving the sample in DMSO-d<sub>6</sub>. We found all samples to be bod-deficient. Together with the formation of NiCl<sub>2</sub>(bod)<sub>2</sub> in preference to NiCl<sub>2</sub>(bod), this suggests that the more electron deficient bod does not coordinate as readily as the btd. This is further borne out by the lack of reported metal complexes containing bod as a ligand in the CSD. We found that the  $x_{\text{NMR}} = 0.31 \ (x_{\text{t}} = 0.75)$  solid-state sample was poorly crystalline and impure, so has not been further analysed. We thus focussed on samples with  $x_t \le 0.75$  for solution state sample and  $x_t \leq 0.50$  for solid state samples.

Comparison of the Pawley derived unit cell volume and lattice parameters with the composition determined from NMR shows linear Vegard's law-type behaviour. The interlayer spacing, c, expands on incorporation of bod, with the M-L-M distance, b, in turn shortening. The contraction along b can be explained by the shorter N-N distance in bod than btd, which would predict  $b_{\rm btd} - b_{\rm bod} = 0.40$  Å, in quantitative agreement with the fitted value of 0.391(7) Å. 19,20 The significant interlayer expansion cannot be easily rationalised by differences in the size between btd and bod, but seem rather to reflect small differences in the chain tilting angles, though might arise from weaker vdW forces for bod than btd. The near constant a axis suggests that the NiCl<sub>2</sub> chain is unperturbed by the differences in Ni-N bonding and that changes in intermolecular forces between bod and btd are not a driving factor. We find no evidence of superlattice reflections indicative of long-range ordering of the bod and btd ligands or structured diffuse scattering from local ordering, though this may be challenging to detect.<sup>21</sup> Having developed this solid-solution series, we then investigated their mechanical compressibility and magnetic properties.

We measured the compressibilities using high pressure synchrotron X-ray powder diffraction (HP-PXRD) at the I15 beamline of Diamond Light Source, using a hydraulic pressure cell to obtain the fine pressure resolution required [Fig. 3].<sup>22</sup> This cell allows measurements from ambient to 0.4 GPa with pressure increments of  $\Delta P = 0.02$  GPa. We used silicone oil AP-100 as a pressure transmitting medium, 23 which should be hydrostatic and non-penetrating at these pressures. We investigated here the doped samples synthesised directly using solid-state synthesis because they were more crystalline. The lattice parameters were refined using Pawley refinement. A limited number of impurity peaks were identified and fitted using additional structure free peaks [Section S2.2, Fig. S6-S14, Tables S2-S4, ESI†].

We found no evidence of pressure-induced framework degradation or phase transitions up to 0.4 GPa. The bulk compressibility, Bo, was fitted using the second-order Birch-Murnaghan equation of state. <sup>25,26</sup> Pure NiCl<sub>2</sub>(btd) has  $B_0 = 18.7(3)$  GPa, with

(a) 0.0 -0.2 -0.4  $\omega^{1}$  -0.6 -0.8

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0.0

**Fig. 3** (a) Linear strain  $(\varepsilon_L)$  along each principal axis determined through HP-PXRD for solid-state NiCl<sub>2</sub>(btd)<sub>1-x</sub>(bod)<sub>x</sub>, with compressibility fit by an empirical equation of state.<sup>24</sup> (b) Linear compressibility indicatrix at P = 0.2 GPa for x = 0.22 (solid) with crystal structure in similar orientation.

0.2

P (GPa)

0.3

the two doped samples both slightly stiffer: x = 0.10 has  $B_0 = 20.6(3)$  GPa and x = 0.22 has  $B_0 = 19.96(13)$  GPa [Fig. S5, ESI†]. These values are comparable to those reported for other Ni(II) vdW materials, *e.g.* Ni(NCS)<sub>2</sub>  $B_0 = 17.0(2)$  GPa,<sup>27</sup> and NiI<sub>2</sub>  $B_0 = 27.7(9)$  GPa,<sup>28</sup> and stiffer than ZnCl<sub>2</sub>(3,5-dichloropyridine)<sub>2</sub>,  $B_0 = 14.52(8)$  GPa, which contains 1D ZnCl<sub>2</sub> chains.<sup>29</sup>

Our HP-XRD measurements probe not only the bulk modulus, but also how compressibility varies with direction. The principal compressibilities approximately coincide with the crystallographic axes, although in a monoclinic system the principal strains will not lie in general along the unit cell axes. The compressibility is largest along the interlayer direction,  $X_1$  $(\sim c) K_1 = 27.3(3) \text{ TPa}^{-1}$ .  $X_2 (\sim a)$  is next stiffest, corresponding to the Ni-Cl-Ni chain direction,  $K_2 = 14.8(4)$  TPa<sup>-1</sup>, with the stiffest direction being the  $X_3$  (b) along the Ni-N-(O/S)-N-Ni bonds direction,  $K_3 = 7.7(4) \text{ TPa}^{-1}$  [Fig. 3 and Table S1, ESI†]. The large compressibility normal to the vdW layers is typical, e.g., Ni(NCS)<sub>2</sub>  $K_{\text{vdW}} = 32.5(2) \text{ TPa}^{-1}$ . As inorganic materials tend to be less compressible,  $X_2$ , with purely inorganic connectivity, might be expected to be the stiffest, but in fact it is nearly twice as soft as  $X_3$ , with purely metal-organic connectivity. This is likely because reducing  $X_3$  corresponds to bond compression, whereas the X<sub>2</sub> direction corresponds to bending of the Ni-Cl-Ni angle, although DFT calculations suggest potentially significant  $\pi$ - $\pi$ -interactions along  $X_2$ . This trend is consistent with previous investigations of metal organic materials, e.g., [CuCl(pyrazine)<sub>2</sub>]BF<sub>4</sub> where the Cu-pyrazine-Cu plane is significantly stiffer than the Cu-Cl-Cu chain, 30 and the plastically deforming ZnCl<sub>2</sub>(3,5-dichloropyridine)<sub>2</sub>, where the ZnCl<sub>2</sub> chains are as soft as the vdW directions ( $K_{\rm ZnCl_2} \approx 23~{\rm TPa}^{-1}$ ).

The changes in compressibility on doping are also anisotropic. The interlayer direction becomes notably stiffer, with compressibility dropping to 24.7(4) TPa<sup>-1</sup> (x = 0.10) and 24.70(11) TPa<sup>-1</sup> (x = 0.20). Within the plane, the inorganic  $X_2$  axis becomes slightly stiffer, 12.6(2) TPa<sup>-1</sup> (x = 0.10) and 14.1(3) TPa<sup>-1</sup> (x = 0.20), whereas the organic  $X_3$  axis in fact softens, 8.14(16) TPa<sup>-1</sup> (x = 0.10) and 8.9(4) TPa<sup>-1</sup> (x = 0.20) [Table S1, ESI†]. Organic substitution thus subtly modifies the compressibility of MOMs, as found for MOFs,<sup>31</sup> and hence can change the efficacy of strain tuning, whether in bulk or on surface.<sup>32,33</sup>

We also investigated the magnetic properties of these MOMs for all five solid-solution samples and NiCl<sub>2</sub>(btd) [Fig. 4 and Section S3, Fig. S21-S24, Table S7, ESI†]. Our previous work showed NiCl<sub>2</sub>(btd) is a canted (weak) ferromagnet due to the noncollinear easy-axes of the paramagnetic Ni<sup>2+</sup> ions with a canting angle of 9.1(4)°.16 All our new samples have similar properties: canted magnets with magnetic ordering temperatures,  $T_c = 17(1)$  K, and substantial hysteresis [Fig. 4(a), (b), (d) and Table S7, ESI $\dagger$ ]. Doping does not greatly affect  $T_c$ , despite the substitution of S for O occurring along the superexchange pathway [Fig. 4(c) and Fig. S22, Table S7, ESI†]. Contrastingly, the more bod added, the softer the magnet, with the coercive field,  $H_c$ , decreasing by 60% on doping with 33% bod (i.e. x =0.33). The reduction in  $H_c$  likely arises as bod has both weaker spin-orbit coupling and ligand field, reducing the single-ion anisotropy. It is also possible that the slight differences in tilt angles between NiCl2 chains induced by ligand geometry change the degree of canting, though this is not clearly observed, and the changes in  $H_c$  are much larger than predicted by geometry alone.

The observed effect of isovalent substitution on magnetic function is consistent with previous studies: replacing S with Se does not produce large changes in  $T_c$  with a 5% reduction for NiPS<sub>3</sub>,<sup>34</sup> and a 30% increase (1.5 K) for Co(NCS)<sub>2</sub>(pyridine)<sub>2</sub>;<sup>35</sup> but does switch the anisotropy from easy-plane in NiPS<sub>3</sub> to easy-axis in NiPSe<sub>3</sub>.<sup>34</sup> We found larger changes in magnetic properties than in the layered ZIFs, where ligand substituents had

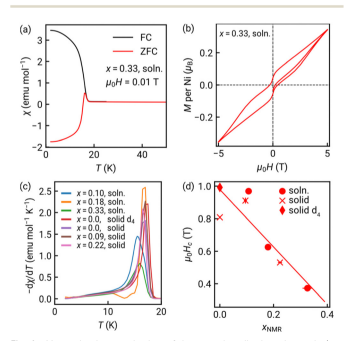


Fig. 4 Magnetic characterisation of the most heavily doped sample (x=0.33, solution). (a) Susceptibility as a function of temperature  $\chi(T)$  for samples cooled in magnetic field (FC) and in zero field (ZFC). Negative magnetisation due to small remnant field in magnet. (b) Isothermal magnetisation, M(H), at T=2 K. The variation in magnetic properties with x, (c) ordering temperature  $T_c$  with x, shown by the peak in  $\frac{\mathrm{d}\chi}{\mathrm{d}T}$  and (d) variation in coercive field  $H_c$ . Linear fit to data shown. Circles indicate solid-state samples (solid), crosses solution sample (soln.), and diamonds solid-state deuterated sample (solid  $\mathrm{d}_d$ ).

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relatively small effects on both  $T_c$  and superexchange. <sup>10,36</sup> The change in coercivity is much larger than previously observed on isovalent substitution in other van der Waals magnets. Pressure can tune noncollinearity in MOMs, <sup>37</sup> and so the combination of mechanical and magnetic tunability we demonstrate suggests that doping will be an effective method to modulate strain switchability.

In conclusion, we demonstrate that ligand solid-solutions achieve fine control over both the mechanical and magnetic properties of vdW MOMs. We report the synthesis of  $\operatorname{NiCl_2(btd)_{1-x}(bod)_x}$  with an approximately linear dependence of the lattice parameters on ligand substitution. The btd ligand is preferentially incorporated into the structure, likely as it is more electron rich. Investigation of the mechanical properties using HP-PXRD showed that bod stiffens the framework, primarily due to a reduction in interlayer compressibility, as the layers themselves become slightly more compressible. The canted ferromagnetism is retained on doping but there is significant reduction (up to 60%) in coercive field. These results demonstrate that functionalisation of organic ligands can be a valuable way to tune both the magnetic function and pressure-responsiveness of van der Waals metal–organic magnets.

E. M., S. L. and J. P.: synthesis. E. M., J. P., S. M. K., S. H., G. K. and D. D.: HP-PXRD measurements. E. M. and S. L.: PXRD and NMR measurements. E. M., S. L. and M. J. C.: NMR and PXRD analysis. E. M., J. P. and M. J. C.: magnetic measurements. E. M. and M. J. C.: magnetic analysis. E. M., S. L. and S. P. A.: SC-XRD measurement and analysis. J. P. T.: electron diffraction measurements. J. P. T. and M. J. C.: electron diffraction analysis. M. J. C. wrote the paper with input from all other authors.

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## Data availability

Data for this article with analysis scripts available at doi: https://doi.org/10.17639/nott.7454. Crystallographic data has

been deposited at the CCDC (https://www.ccdc.cam.ac.uk/structures/) with reference numbers 2373766, 2373767 and 2377919 †

#### Conflicts of interest

There are no conflicts to declare.

#### Notes and references

- 1 M. Eddaoudi, et al., Science, 2002, 295, 469-472.
- 2 X. Kong, et al., Science, 2013, 341, 882-885.
- 3 A. D. Burrows, et al., Chem. Commun., 2011, 47, 3380-3382.
- 4 S. Horike, et al., Chem. Sci., 2012, 3, 116-120.
- 5 C.-C. Cao, et al., J. Am. Chem. Soc., 2019, 141, 2589-2593.
- 6 P. Vervoorts, et al., ACS Mater. Lett., 2021, 3, 1635-1651.
- 7 A. E. Thorarinsdottir and T. D. Harris, Chem. Rev., 2020, 120, 8716–8789.
- 8 L. S. Xie, et al., Chem. Rev., 2020, 120, 8536-8580.
- 9 A. Helal, et al., Natl. Sci. Rev., 2017, 4, 296-298.
- 10 J. Lopez-Cabrelles, et al., Chem. Sci., 2022, 13, 842-847.
- 11 J. Song, et al., Angew. Chem., Int. Ed., 2022, 61, e202117565.
- 12 T. Li, et al., Nat. Mater., 2019, 1-6.
- 13 F. Aribot, et al., ChemRxiv, 2024, preprint, DOI: 10.26434/chemrxiv-2024-rlsf8.
- 14 S. Mondal, et al., Phys. Rev. B: Condens. Matter Mater. Phys., 2019, 99, 180407.
- 15 C. A. Occhialini, et al., arXiv, 2023, preprint, arXiv:2306.11720, DOI: 10.48550/arXiv.2306.11720.
- 16 J. Pitcairn, et al., J. Am. Chem. Soc., 2024, 146, 19146-19159.
- 17 G. S. Papaefstathiou, et al., J. Solid State Chem., 2001, 159, 371-378.
- 18 B. Morosin, Acta Crystallogr., 1967, 23, 630-634.
- 19 T. Suzuki, et al., J. Org. Chem., 2001, 66, 8954-8960.
- 20 M. R. Ams, et al., Chem. Eur. J., 2019, 25, 323-333.
- 21 M. L. Robinson, et al., Inorg. Chem., 2020, 59, 3026-3033.
- 22 N. J. Brooks, et al., Rev. Sci. Instrum., 2010, 81, 064103.
- 23 S. Dissegna, et al., J. Am. Chem. Soc., 2018, 140, 11581-11584.
- 24 A. L. Goodwin, et al., Proc. Natl. Acad. Sci. U. S. A., 2008, 105, 18708–18713.
- 25 F. Birch, Phys. Rev., 1947, 71, 809-824.
- 26 M. Lertkiattrakul, et al., J. Open Source Softw., 2023, 8, 5556.
- 27 M. Geers, et al., Phys. Rev. B, 2023, 108, 144439.
- 28 M. P. Pasternak, et al., Phys. Rev. Lett., 1990, 65, 790-793.
- 29 X. Liu, et al., Nat. Commun., 2021, 12, 3871.
- 30 R. Scatena, et al., Inorg. Chem., 2020, 59, 10091-10098.
- 31 L. R. Redfern, et al., J. Am. Chem. Soc., 2019, 141, 4365-4371.
- 32 D. L. Esteras, et al., Nano Lett., 2022, 22, 8771–8778.
- 33 J. Cenker, et al., Nat. Nanotechnol., 2022, 17, 256–261.
- 34 R. Basnet, et al., Phys. Rev. Res., 2022, 4, 023256.
- 35 T. Neumann, et al., Chem. Commun., 2019, 55, 2652-2655.
- 36 J. López-Cabrelles, et al., J. Am. Chem. Soc., 2021, 143, 18502-18510.
- 37 I. E. Collings, et al., Phys. Chem. Chem. Phys., 2018, 20, 24465-24476.