(23)Na multiple-quantum MAS NMR of the perovskites NaNbO(3) and NaTaO(3).
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The distorted perovskites NaTaO$_3$ and NaNbO$_3$ have been studied using $^{23}$Na multiple-quantum (MQ) MAS NMR. NaTaO$_3$ was prepared by high temperature solid state synthesis and the NMR spectra are consistent with the expected room temperature structure of the material (space group $P$mm), with a single crystallographic sodium site. Two samples of NaNbO$_3$ were studied. The first, a commercially available sample which was annealed at 900 °C, showed two crystallographic sodium sites, as expected for the room temperature structure of the material (space group $P$bcm). The second sample, prepared by a low temperature hydrothermal method, showed the presence of four sodium sites, two of which match the expected room temperature structure and the second pair, another polymorph of the material (space group $P2_1$). This is consistent with powder X-ray diffraction data which showed weak extra peaks which can be accounted for by the presence of this second polymorph. Density functional theory (DFT) calculations support our conclusions, and aid assignment of the NMR spectra. Finally, we discuss the measured NMR parameters in relation to other studies of sodium in high coordination sites in the solid state.

Introduction

The use of $^{23}$Na NMR for the structural characterisation of the local environment of sodium in the solid state has attracted considerable attention recently for a wide range of chemical systems. Particularly well-studied have been silicate minerals where sodium is found in diverse and often highly unsymmetric coordination environments. Owing to the variety of sodium chemical environments, it can be difficult to relate directly experimental NMR parameters (such as isotropic chemical shift) with coordination geometry (average bond distances and coordination numbers). In addition, $^{23}$Na has spin quantum number $I = 3/2$, with spectra broadened by the quadrupolar interaction, even under MAS. This hinders the determination of accurate quadrupole and chemical shift parameters, particularly when a number of crystallographically distinct species are present. However, the development of techniques such as multiple-quantum (MQ) MAS and satellite-transition (ST) MAS, capable of removing completely both first- and second-order quadrupolar broadening, have enabled high-resolution $^{23}$Na NMR spectra to be recorded routinely and NMR parameters to be extracted accurately and easily.

Despite the fairly small chemical shift range of $^{23}$Na, some general trends with structure have been observed, particularly within groups of similar materials. For example, a number of correlations between isotropic chemical shift ($\delta_{CS}$) and Na–O distances and coordination number have been observed, particularly for silicates and fluorides, and whilst they are not identical, it is clear that sodium atoms with a larger number of oxygen or fluorine near neighbours (usually, therefore, with a higher average bond distance) exhibit a lower chemical shift. The significant progress in the development of ab initio calculations of NMR parameters over the past few years has also facilitated the study of the variation of NMR parameters with structural changes. Many early calculations utilised a cluster approach, ultimately limited in accuracy (and indeed feasibility) by cluster size. Twenty years ago, Schwarz and co-authors showed that the linear augmented plane-wave (LAPW) method was able to predict with high precision electric field gradients in solids. Recent developments of the projector augmented wave formalism (PAW) for the calculation of electric field gradients, and its extension, the gauge including projector augmented wave formalism (GIPAW), enabling calculation of both electric field gradients and chemical shielding tensors, now allow accurate calculations to be performed for periodic solids and correlations investigated.

In this work we use MQMAS to acquire high-resolution spectra of the oxides NaNbO$_3$ and NaTaO$_3$, and employ density functional theory (DFT) calculations to aid assignment and interpretation of our spectra. The two materials have been characterised structurally by a number of groups and there exists in the literature several reports of...
crystallographic studies of each material.\textsuperscript{17–23} Both exhibit a rich polymorphism over wide ranges of temperature, but all structures observed are the result of displacive transitions, involving octahedral tilts or rotations within the ideal, cubic ABO\textsubscript{3} perovskite structure. Here, the sodium resides on the (ideally) twelve-coordinate A site, and the Group 5 metal occupies the octahedral B site. At room temperature, NaNbO\textsubscript{3} is usually reported to possess an orthorhombic unit cell with $a = 5.506$ Å, $b = 5.566$ Å and $c = 15.52$ Å in space group Pbnm\textsuperscript{17} (some authors have used the alternative setting Pbma) in which two crystallographic sodium sites are found, as in Fig. 1b. A powder neutron diffraction experiment by Darlington and Knight indicated that the symmetry of NaNbO\textsubscript{3} at room temperature might be lower than orthorhombic;\textsuperscript{21} although another group have subsequently found no evidence for this in a similar study.\textsuperscript{23} NaNbO\textsubscript{3} has been reported to adopt a structure with an orthorhombic unit cell at room temperature with $a = 5.4768$ Å, $b = 5.5212$ Å and $c = 7.7890$ Å in space group Pbnm, but with a smaller volume unit cell than NaNbO\textsubscript{2}, which includes only one crystallographic sodium site,\textsuperscript{22} as in Fig. 1a. Only a few previous \textsuperscript{23}Na NMR studies of NaNbO\textsubscript{3} have been described. For example, an early study by Wolf et al.,\textsuperscript{24} performed by field sweeping with $\nu_0 = 10$ MHz, was able to determine the presence of two distinct Na sites from the position of singularities in the NMR spectrum, whilst \textsuperscript{23}Na NMR of NaNbO\textsubscript{2} was mentioned, but not shown directly, in the work of Haase et al.\textsuperscript{25} Aside from the interest in obtaining high-resolution \textsuperscript{23}Na MQMAS spectra of the two perovskites, where Na has an unusually high coordination number, the complexity of the phase diagrams of NaNbO\textsubscript{3} and NaTaO\textsubscript{3}, with the possible existence of various polymorphs even at room temperature, suggest \textsuperscript{23}Na NMR may be a useful tool for sample characterisation.

### Experimental

#### Sample preparation

NaTaO\textsubscript{3} was prepared using the method of Kennedy et al.\textsuperscript{22} Stoichiometric amounts of dry Na\textsubscript{2}CO\textsubscript{3} (BDH, 99.9\%) and Ta\textsubscript{2}O\textsubscript{5} (Aldrich 99\%) were finely ground and fired at 700, 800 and 900 °C for 24 h at each temperature with intermittent regrounding, followed by a final firing at 1150 °C. Powder X-ray diffraction (recorded using a Bruker D8 diffractometer operating with Cu K\textalpha radiation, average $\lambda = 1.5418$ Å) confirmed that the solid-state reaction was complete and all reflections could be indexed using the 300 K structural model of Kennedy et al.\textsuperscript{22}

A first sample of NaNbO\textsubscript{3} was supplied by Aldrich (100 mesh, no purity stated) and its crystallinity improved by annealing at 900 °C for three days. Powder X-ray diffraction data from this material were consistent with the room temperature structural model deduced by Sakowski-Cowley et al.\textsuperscript{17} A second sample of NaNbO\textsubscript{3} was prepared using a hydrothermal method, based on that of Goh et al.,\textsuperscript{26} 0.67 g of Nb\textsubscript{2}O\textsubscript{5} (Aldrich 99.99\%) were stirred into 10 ml of 8.4 M NaOH solution and heated in a 23 ml Teflon-lined, stainless-steel autoclave at 200 °C for 24 h. The white solid was recovered by suction filtration, washed copiously with deionised water and dried at 100 °C in air overnight. Powder X-ray diffraction of the second sample gave data in broad agreement with the expected orthorhombic model of Sakowski-Cowley et al.,\textsuperscript{17} but weak extra reflections were also observed, as shown in Fig. 2. After searching the Inorganic Crystal Structure Database (ICSD), we discovered that the only possible polymorph of NaNbO\textsubscript{3} that matched these extra peaks was one with space group P2\textsubscript{1}1ma reported by Shuvaeva et al.,\textsuperscript{27} also represented on Fig. 1c. Simulating the powder X-ray pattern as a 50 : 50 mixture of the Pbnm and P2\textsubscript{1}1ma polymorphs gave a good fit to the observed powder diffraction data. We will discuss this conclusion further, in the light of the NMR results below.

#### NMR spectroscopy

\textsuperscript{23}Na MAS NMR spectra were obtained at Larmor frequencies of 52.9, 105.8, 132.2 or 158.7 MHz on Bruker Avance 200, Chemagnetics Infinity 400, Varian Infinity Plus 500 and Bruker Avance 600 spectrometers, respectively. Powdered samples were packed inside either 4-mm or 2.5-mm MAS rotors and rotated at speeds between 10–25 kHz. Spectra are referenced to 1 M NaCl (aq) through a secondary reference of NaCl (s) at 7.3 ppm. Two-dimensional triple-quantum MAS spectra were recorded using the phase-modulated split-t\textsubscript{1} shifted-echo sequence given in Fig. 12a of ref. 28.
radiofrequency field strengths employed were between 80–150 kHz for the triple-quantum excitation and conversion pulses and 10–20 kHz for the final inversion pulse. All spectra were acquired at room temperature (B18C). Any further experimental details may be found in the figure captions.

Calculations

DFT calculations were carried out using the WIEN2k package,29 and in some cases the NMR-CASTEP code.30 WIEN2k is a full potential all electron method that utilizes the L/APW + lo approach.31 The exchange and correlation potentials were calculated using the generalized gradient approximation (GGA) in the parameterization of Perdew–Burke–Ernzerhof.32 Atomic radii of 1.8, 1.8, 1.4, and 2.5 bohr were used for Nb, Ta, O, and Na spheres, respectively. Charge self-consistency was obtained from 9 and 48 irreducible k points for the calculation of NaNbO3 structures Pbcm and P21ma, respectively. The Brillouin zone integration was performed with 18 irreducible k points for both NaTaO3 structures. The full electric field gradient tensor was calculated from the total self-consistent charge density in the crystal without further approximation. As all electrons are taken into account, the charge redistribution is automatically achieved in the self-consistent field procedure, and thus no additional Sternheimer antishielding factors are required. A 23Na quadrupole moment of 104 mb was employed.33 As the sign of CQ is, in most cases, not able to be determined easily directly from a simple MAS spectrum, absolute values of CQ have been considered.

Both quadrupolar and chemical shift parameters were also calculated using the NMR-CASTEP30 code, which calculates the chemical shieldings and electric field gradients within periodic boundary conditions and the pseudopotential approximation using the GIPAW13 and PAW11–12 methods. Ultrasoft Vanderbilt pseudopotentials were used, with 2s and 2p valence for O, 2s, 2p and 3s valence for Na and 4s, 4p, 5s and 4d valence for Nb. Calculations were performed with a cut-off energy of 689 eV and with 0.05 k points Å3 using GGA (PBE32). Geometry optimisations were also performed using this code, with the NMR parameters for the optimised structures then calculated using WIEN2k and/or NMR-CASTEP.

In all calculations (WIEN2k and NMR-CASTEP) a unit cell of the material was considered (8 formula units for NaNbO3, Pbcm and 4 formula units for NaNbO3, P21ma, and both polymorphs of NaTaO3), with the experimental crystallographic parameters (unit cell, atomic positions and space group) as shown in Fig. 1, unless specified.

Results

The 23Na MAS NMR spectra of NaTaO3 recorded at a range of field strengths, B0, are shown in Fig. 3. All spectra exhibit a single second-order broadened lineshape, with a width (in Hz) proportional to 1/B0. The isotropic chemical shift ($\delta_{CS}$) and the quadrupolar coupling, (CQ = $e^2Qq/h$) and asymmetry ($\eta$) may be determined by computer fitting of this lineshape, using the dmfit program,34 yielding the values given in Table 1. A comparison of the experimental 23Na spectra with those simulated using the fitted parameters is also shown in Fig. 3. (It should be noted that this method of fitting becomes significantly more difficult as the number of crystallographically-distinct species increases owing to overlap of the quadrupolar broadened lineshapes.) Two-dimensional multiple-quantum MAS spectra (shown for B0 = 9.4 T in Fig. 4) also reveal a single ridge, parallel with the $\delta_{CS}$ axis. The isotropic (or high-resolution) spectrum, obtained from a projection onto $\delta_{CS}$, contains a single, sharp resonance. For MQMAS spectra in general, it is possible to extract information on the quadrupolar and chemical shift interactions from the spectra using a
variety of methods. The position of the centre of gravity of the ridge lineshape ($\delta_1$, $\delta_2$) enables determination of $\delta_{CS}$ and $P_Q$, the quadrupolar product, given by $C_Q (1 + \eta^2/3)^{1/2}$. Alternatively, $C_Q$, $\eta$ and $\delta_{CS}$ may be obtained by computer fitting of a cross-section extracted along the ridge lineshape, parallel to $\delta_2$, in a similar manner to the one-dimensional spectrum, although this lineshape may sometimes be distorted as a result of non-uniform excitation and conversion of triple-quantum coherences, as indeed is the case for the spectrum shown in Fig. 4. Finally, it is also possible to determine $\delta_{CS}$ and $P_Q$ from the variation of the $\delta_1$ position of a ridge with $B_0$ field strength. For example, if $\delta_1$ (in ppm) is plotted against $(1/\nu_0^2)$, where $\nu_0$ is the Larmor frequency in Hz, the result will be (for $I = 3/2$) a straight line with gradient $(31250 P_Q^2)$ and intercept $(17 \delta_{CS}/8)$. The values of quadrupolar and chemical shift parameters determined from the MQMAS spectra are also given in Table 1.

Table 1

<table>
<thead>
<tr>
<th></th>
<th>$\delta_{CS}$ (ppm)</th>
<th>$C_Q$/MHz</th>
<th>$\eta$</th>
<th>$P_Q$/MHz</th>
</tr>
</thead>
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<tr>
<td>MAS</td>
<td>$-3.0 \pm 1.0$</td>
<td>$1.3 \pm 0.1$</td>
<td>$0.9 \pm 0.1$</td>
<td>$1.5^*$</td>
</tr>
<tr>
<td>MQMAS</td>
<td>$-2.5 \pm 1.0$</td>
<td>$1.3 \pm 0.1$</td>
<td>$0.95 \pm 0.1$</td>
<td>$1.6 \pm 0.1$</td>
</tr>
</tbody>
</table>

* Calculated from $C_Q (1 + \eta^2/3)^{1/2}$.

Fig. 4 Two-dimensional $^{23}$Na triple-quantum MAS spectrum of NaTaO$_3$, recorded using a phase-modulated split-$t_1$ shifted echo pulse sequence, with $B_0 = 9.4$ T. The spectrum is the result of averaging 192 transients with a recycle interval of 2 s, for each of 128 $t_1$ increments of 50 $\mu$s. The MAS rate was 12 kHz. The isotropic projection (onto the $\delta_1$ axis) and cross-section parallel to $\delta_2$ are also shown. All chemical shifts are shown relative to 1 M NaCl (aq).

The parameters determined from all three field strengths and from different types of spectra are in excellent agreement with each other. The values of the quadrupolar coupling constant and asymmetry are also in good agreement with those calculated by DFT using the WIEN2k package for the room temperature Pbnm polymorph, and given in Table 2. However, at all field strengths there are very small discrepancies between the experimental and simulated lineshapes (most noticeable on the exact positions and heights of the distinct shoulders on each lineshape), with the experimental MAS and MQMAS spectra appearing to exhibit a small additional broadening. There are a few possible explanations for such broadening. Perhaps the simplest is that of some extent of distortion within the system. This would produce distributions of both quadrupolar and chemical shift parameters which would additionally broaden the spectra. If such distributions are present they must be fairly small as no significant additional broadening is observed in the two-dimensional MQMAS spectra. An alternative explanation could involve the presence of a higher-order interaction which is unable to be averaged by MAS, such as a second-order quadrupolar–dipolar cross term, arising from dipolar coupling of Na to a nearby quadrupolar $^{181}$Ta ($I = 7/2$) nucleus. Usually such interactions are very small, but the large nuclear quadrupolar

Table 2

<table>
<thead>
<tr>
<th>Space group</th>
<th>$\delta_{CS}$ (ppm)</th>
<th>$C_Q$/MHz</th>
<th>$\eta$</th>
<th>$P_Q$/MHz</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pbnm (Na1)</td>
<td>—</td>
<td>1.29</td>
<td>0.95</td>
<td>1.48$^*$</td>
</tr>
<tr>
<td>Cmcm (Na1)</td>
<td>—</td>
<td>1.34</td>
<td>0.44</td>
<td>1.38$^*$</td>
</tr>
<tr>
<td>Cmcm (Na2)</td>
<td>—</td>
<td>0.63</td>
<td>0.15</td>
<td>0.63$^*$</td>
</tr>
</tbody>
</table>

* Calculated from $C_Q (1 + \eta^2/3)^{1/2}$.
moment of $^{181}$Ta (32 times greater than $^{23}$Na) may make this effect small but significant. One final possibility is that the broadening may be due to the presence of a small amount of the high-temperature $Cmcm$ phase, undetected by X-ray diffraction. This phase has been reported as existing over the temperature range 723–843 K, but has been found as a minor impurity in samples at room temperature. The $^{23}$Na quadrupolar parameters for the two distinct Na species in this phase, calculated by WIEN2k are given in Table 2. It was not possible to detect unambiguously the presence of these resonances in the two-dimensional $^{23}$Na spectra, however, although signals with very low intensity (<2% of the maximum and below the contour levels displayed in Fig. 4) were observed.

Fig. 5 shows $^{23}$Na conventional MAS and two-dimensional triple-quantum MAS NMR spectra of NaNbO$_3$ (purchased from Aldrich) at field strengths $B_0$ of 4.7 and 11.7 T. In each case, two distinct resonances, overlapped in the MAS spectrum, are resolved in the MQMAS spectrum. Initial spectra acquired on this sample as purchased exhibited additional broadening as a result of poor crystallinity, but this was improved by annealing the material. The differing widths of the two resonances in Fig. 5 reveal that the two species have very different quadrupolar couplings. This is confirmed in Table 3, which gives the quadrupolar and chemical shift parameters extracted (using the variety of methods described above) from the MAS and MQMAS spectra for the two distinct Na species. This difference in quadrupolar coupling constant (2.1 and 1.0 MHz, respectively, for Na(1) and Na(2)), is responsible for the differing relative intensities observed in the MQMAS spectra. As described previously, the excitation of triple-quantum coherences is non-uniform as the magnitude of the quadrupolar coupling changes. This manifests itself both as lineshape distortions (reflecting the dependence of the quadrupolar interaction upon crystallite orientation) and in non-quantitative signal intensities, as observed in Fig. 5, particularly at the lower $B_0$ field strength.

The quadrupolar coupling constants obtained for NaNbO$_3$ are in excellent agreement with those obtained from DFT calculations using WIEN2k (room temperature $Pbcm$ polymorph), given in Table 4. However, the agreement of the asymmetry parameter, whilst reasonably good for Na(2), is not in as good agreement for Na(1), where the calculated parameter appears significantly higher than that observed experimentally. This disagreement was also noted in the early work of Wolf et al., where point charge calculations of the Na electric field gradients resulted in good agreement for $C_Q$, but for Na(1), $\eta$ ranged between 0.6 and 0.72, in contrast to the experimental value of $\sim 0$. Electric field gradient results yielded by NMR-CASTEP (not shown) were in general agreement with those from WIEN2k and also showed a non-zero $\eta$ for Na(1). However, DFT calculations revealed a force of 2.4 eV Å$^{-1}$ for each Nb atom (much larger than that usually expected $\sim 0.1$ eV Å$^{-1}$), suggesting that its position, as determined by X-ray diffraction, is not optimal. Therefore, a full geometry optimisation was performed (using NMR-CASTEPE), allowing the position of all atoms to change, and resulted in a much lower Na(1) $\eta$ of $\sim 0.01$ using WIEN2k, but a increase in the $C_Q$ was also observed (up to 2.5 MHz). Geometry optimisation allowing only the position of the Nb to be altered did not result in any significant change to the Na NMR parameters determined by WIEN2k from those

### Table 3

| $^{23}$Na NMR parameters for NaNbO$_3$ (Aldrich) from the MAS and MQMAS spectra in Fig. 5 |
|-----------------|-----------------|-----------------|-----------------|
| $\delta_{CS}$ (ppm) | $C_Q$/MHz | $\eta$ | $P_Q$/MHz |
| Na(1) | $-0.5 \pm 1.0$ | $2.1 \pm 0.1$ | $0.0 \pm 0.1$ | $2.1 \pm 0.1$ |
| Na(2) | $-4.5 \pm 1.0$ | $1.0 \pm 0.1$ | $0.9 \pm 0.2$ | $1.2 \pm 0.1$ |

### Table 4

| Calculated $^{23}$Na NMR parameters for NaNbO$_3$ polymorphs using WIEN2k |
|-----------------|-----------------|-----------------|-----------------|
| Space group | $\delta_{CS}$ (ppm) | $C_Q$/MHz | $\eta$ | $P_Q$/MHz |
| $Pbcm$ (Na1) | $-0.5 \pm 1.0$ | $2.1 \pm 0.1$ | $0.0 \pm 0.1$ | $2.1 \pm 0.1$ |
| $Pbcm$ (Na2) | $-4.5 \pm 1.0$ | $1.0 \pm 0.1$ | $0.9 \pm 0.2$ | $1.2 \pm 0.1$ |

* Calculated from $C_Q(1 + \eta^2)^3/12$. |
obtained initially. These results were also confirmed using NMR-CASTEP. Given the debate in the literature\textsuperscript{17–23,27,36} over the exact symmetry of the NaNbO\textsubscript{3} structure, perhaps the most likely explanation is a disordering of the Nb position within the NbO\textsubscript{6} octahedra, either time-dependent (dynamic) or static, with only an average value observed by diffraction. Alternatively, the structure may be of a lower symmetry than previously thought. In this respect it should be noted that whilst well-defined \textsuperscript{23}Na quadrupolar-broadened lineshapes are obtained, \textsuperscript{93}Nb MAS and MQMAS spectra of NaNbO\textsubscript{3} show a distinctly broader lineshape (with $C_Q \approx 20$ MHz) but with few characteristic features of second-order broadening, perhaps confirming the presence of Nb disorder.\textsuperscript{37,38} It should also be noted that it is difficult to rule out completely any effects of Na ion motion on the spectra, although these are expected to be small.

Fig. 6 shows similar \textsuperscript{23}Na MAS and MQMAS spectra (at $B_0$ field strengths of 4.7, 9.4, 11.7 and 14.1 T) for a sample of NaNbO\textsubscript{3} prepared using a hydrothermal method,\textsuperscript{26} as described in the experimental section. The MAS spectra look similar to those obtained for the Aldrich sample, although the sharper singularities reflect the better crystallinity of this hydrothermal sample, despite the annealing of the as-purchased material. However, significant differences between the two samples are observed in the MQMAS spectra, with additional resonances present in the spectrum of the hydrothermal material. There is significant overlap of spectral resonances and all cannot easily be resolved at a single $B_0$ field strength. At high $B_0$, the signal at lower $\delta_1$ appears to be comprised of two closely spaced resonances, whilst at higher $\delta_1$, ppm, produces a single (albeit broadened) resonance in the isotropic spectrum. Conversely, at low $B_0$, this resonance is clearly resolved into two distinct peaks, whilst only a single peak is observed at low $\delta_1$. This suggests that the spectrum is composed of four distinct resonances, two with lower $C_Q$, which are resolved at higher $B_0$, owing to their differing chemical shifts, and two with much higher $C_Q$, resolved at lower $B_0$, as a result of differing second-order quadrupolar shifts. That these latter two resonances are coincident in $\delta_1$ at higher $B_0$ can also be observed by considering the cross-sections extracted parallel to the $\delta_2$ axis in each spectrum, shown in Fig. 7. The lineshape at 14.1 T (with

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig6.png}
\caption{\textsuperscript{23}Na MAS NMR of NaNbO\textsubscript{3} synthesised by a hydrothermal method. Conventional MAS spectra, two-dimensional triple-quantum MAS (recorded using a phase-modulated split-$t_1$ shifted echo pulse sequence\textsuperscript{28}) and corresponding isotropic projections are shown for magnetic field strengths, $B_0$, of (a) 4.7, (b) 9.4, (c) 11.7 and (d) 14.1 T. MAS spectra result from the averaging of (a) 480, (b, c, d) 240 transients with a recycle interval of 2 s. Two-dimensional MQMAS spectra result from the averaging of (a) 192, (b) 576, (c) 288 and (d) 192 transients with a recycle interval of 2 s for each of (a) 256, (b) 128, (c) 115 and (d) 192 $t_1$ increments of (a) 33.3, (b) 50, (c) 110.2 and (d) 148.1 $\mu$s. The MAS rate was (a) 8, (b) 12, (c) 16 and (d) 12.5 kHz. All chemical shifts are shown relative to 1 M NaCl (aq).}
\end{figure}


Fig. 7 Cross-sections extracted parallel to the $\delta_2$ axis along the ridge lineshapes in the two-dimensional $^{23}$Na triple-quantum MAS NMR spectra of NaNbO$_3$ synthesised by a hydrothermal method, shown in Fig. 6(a)–(d). All chemical shifts are shown relative to 1 M NaCl (aq).

$\delta_1$ = 4 ppm) appears to result from the overlap of two different quadrupolar-broadened powder pattern lineshapes with distinctly different $\eta$. Although resolved at low $B_0$, the lineshapes are then somewhat distorted as a result of non-uniform excitation of triple-quantum coherences, and it is more difficult to extract accurate values for $\eta$.

The complexity of these spectra, and particularly the considerable peak overlap, does hinder the extraction of accurate quadrupolar and chemical shift parameters both from the position of the centre of gravity of each lineshape and from cross-sections at particular $B_0$ field strengths. It should be noted however, that the $\delta_1$ positions of each resonance remain well defined at all field strengths, even when considerable overlap is present. This relationship of $\delta_1$ with $B_0$ field may be used to determine $\delta_{CS}$ and $P_Q$ (as described above). Values for the quadrupole and chemical shift parameters extracted using this approach (and from the centre of gravity or from cross-sections when possible) are given in Table 5 for the four distinct resonances, labelled 1, 2, 3 and 4 in order of increasing $\delta_1$ shift. A comparison of Fig. 5 and 6 and Tables 3 and 5 reveal that resonances 2 and 3 can be clearly identified as Na(2) and Na(1), respectively of NaNbO$_3$ (Pbcm), observed in the as purchased sample.

Table 5 $^{23}$Na NMR parameters for NaNbO$_3$ (hydrothermal synthesis) from the MAS and MQMAS spectra in Fig. 6 and 7

<table>
<thead>
<tr>
<th>$\delta_{CS}$ (ppm)</th>
<th>$C_Q$/MHz</th>
<th>$\eta$</th>
<th>$P_Q$/MHz</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$-5.0 \pm 2.0$</td>
<td>$1.0 \pm 0.1$</td>
<td>$0.7 \pm 0.2$</td>
</tr>
<tr>
<td>2</td>
<td>$-4.5 \pm 2.0$</td>
<td>$1.0 \pm 0.1$</td>
<td>$0.8 \pm 0.2$</td>
</tr>
<tr>
<td>3</td>
<td>$-1.0 \pm 2.0$</td>
<td>$2.1 \pm 0.1$</td>
<td>$0.0 \pm 0.2$</td>
</tr>
<tr>
<td>4</td>
<td>$-1.5 \pm 2.0$</td>
<td>$2.1 \pm 0.1$</td>
<td>$0.9 \pm 0.2$</td>
</tr>
</tbody>
</table>

Peaks labelled 1 to 4 in order of increasing $\delta_1$ shift.

To examine the possibility of additional resonances originating from the $P21ma$ polymorph, as suggested by the powder diffraction data, we again performed DFT calculations using WIEN2k, the results of which are given in Table 4. The quadrupolar parameters of the two sodiums are very similar to those obtained experimentally, suggesting that our hydrothermal synthesis has indeed resulted in a mixture of the $Pbcm$ and $P21ma$ polymorphs. NMR-CASTEP calculations confirmed the values of these Na electric field gradients and predicted a $\sim 4$ ppm difference in the two chemical shifts, as observed experimentally. The relative intensities of the resonances seen in the 4.7 T spectrum suggest that approximately equal amounts of each polymorph are present, consistent with the powder X-ray diffraction data. The NMR parameters are very similar to those obtained for the two resonances in $Pbcm$ NaNbO$_3$, resulting in considerable spectral overlap in the MAS spectra, and indeed in the MQMAS spectra at particular field strengths.

It should also be noted that although small differences are observed in the $^{93}$Nb MAS spectrum for this material to that for the commercially-available material, the broadened lines observed (as discussed above) prevent the extraction of any detailed information on the number or type of polymorphs present.

Discussion and conclusions

The above results demonstrate the sensitivity of $^{23}$Na NMR to local structure and also highlight the need for high-resolution NMR techniques when studying quadrupolar nuclei. Despite very similar MAS spectra, the two samples of NaNbO$_3$ (as purchased and prepared by hydrothermal synthesis) have very different MQMAS spectra, with the presence of two additional resonances in the latter. It should also be noted that full characterisation of these materials required experiments to be performed at more than one $B_0$ field strength as a result of the differing field dependences of the chemical shift (proportional, in Hz, to $B_0$) and the second-order quadrupolar interaction (proportional, in Hz, to $1/B_0^2$), and the similarity of the species present. The fairly large differences in both quadrupolar coupling and chemical shift of the two Na species in the room temperature $Pbcm$ NaNbO$_3$, resulting from the distortion or tilting of the NbO$_6$ octahedra, also emphasizes the sensitivity of NMR to small changes in the local environment.

The $^{23}$Na NMR study of the sample of NaNbO$_3$ prepared using hydrothermal synthesis provides independent, complementary evidence to the powder diffraction data that the material contains a mixture of two polymorphs of the...
material. In addition to the expected \textit{Pbcm} polymorph we have found convincing evidence that the sample contains a significant proportion of a second polymorph, probably \textit{P21ma} (around 50\% by mass). The previous report of this polymorph found that the phase was produced in the presence of an electric field, inducing small shifts in the positions of two unique oxide ions, resulting in a structure similar to that of \(K_{0.025}Na_{0.975}NbO_3\).\textsuperscript{27} It is most likely that it is our use of a low temperature synthetic route (200 °C, instead of in excess of 1000 °C) that has allowed access to this unusual polymorph. It is interesting to note that studies of the synthesis of the perovskite barium titanate, \(BaTiO_3\), have shown that a cubic polymorph, usually only seen above 100 °C, may be formed using mild hydrothermal conditions.\textsuperscript{39} Previous studies of the hydrothermal synthesis of \(NaNbO_3\) used low-resolution powder X-ray diffraction for phase identification,\textsuperscript{26} and so the presence of any weak reflections due to the second polymorph may have been difficult to observe. This demonstrates the power of combining powder diffraction with solid-state NMR (particularly when using calculations to aid spectral assignment) in the characterisation of polycrystalline samples.

As discussed in the Introduction, several authors have noted a general trend of decreasing isotropic chemical shift with increasing average sodium coordination number (and hence average Na–O distance).\textsuperscript{3–9} Our newly measured values for sodium \(NaTaO_3\) and \(NaNbO_3\) fall between 0 and –6 ppm, consistent with this general trend. Despite the fact that sodium is often found in highly irregular coordination environments, Koller \textit{et al.}\textsuperscript{5} determined that all oxygen atoms in a sphere with a radius of 3.4 Å around sodium contributed to the observed chemical shift of the sodium and could be considered as part of the immediate coordination environment. Both \(NaTaO_3\) and \(NaNbO_3\) have structures that are distortions of the ideal, cubic perovskite structure and the nominally twelve-coordinate A site is not a regular cubooctahedron. For \(NaTaO_3\), the single crystallographic sodium is surrounded by 12 oxygens at distances ranging from 2.419–3.173 Å, with a mean value of 2.764 Å,\textsuperscript{22} whilst for \(NaNbO_3\) (\textit{Pbcm}), Na1 has Na–O distances ranging from 2.387–3.129 Å (mean 2.770 Å), and Na2 has Na–O distances ranging from 2.383–3.171 Å (mean 2.777 Å).\textsuperscript{17} For the \textit{P21ma} polymorph, one sodium has Na–O distances ranging from 2.372–3.148 Å (mean 2.780 Å), and the other, distances ranging from 2.484–3.272 Å (mean 2.784 Å).\textsuperscript{22}

Koller \textit{et al.}\textsuperscript{5} quantified the relationship between chemical shift and local sodium coordination by considering the bond valence sum of each individual oxygen, \(W_i\),\textsuperscript{40} and then introducing a shift parameter, \(A\), for each unique sodium according to the equation:

\[
A = \sum_j \left( W_i / r_j^7 \right)
\]  

Here the sum is performed over all cation(j)–oxygen(i) pairs, and \(n\) is ideally 3. For the materials studied by Koller \textit{et al.},\textsuperscript{5} the majority of which were silicates and phosphates, a linear relationship between isotropic chemical shift and \(A\) was observed. We have found that for \(NaTaO_3\) and \(NaNbO_3\), the measured values of chemical shift do not fit this correlation, and in fact much more negative chemical shifts would be expected if the correlation was extrapolated to the \(A\) values for sodium in these perovskite materials. For example, for \(NaTaO_3\), \(A\) for the unique sodium is 1.28, giving an expected isotropic chemical shift of –55.6 ppm (correcting the chemical shift to be referenced to NaCl (s) at 7.3 ppm as in our measured data). The only transition-metal oxide material that Koller \textit{et al.} studied was \(NaCrO_3\), which contains two unique sodiums, one of which has ten oxygen neighbours at less than 3.4 Å, at an average distance of 2.86 Å (note this is a longer average distance than for the 12 coordinate sodium ions found in the perovskite materials) and was observed to have an isotropic chemical shift of –12.2 ppm. In a separate work, Xue and Stebbins\textsuperscript{4} studied a series of silicates and noted that a correlation of the chemical shift with the average Na–O interatomic distance is often more applicable than the correlation of Koller \textit{et al}. It was noted in this work that in many simple systems the determination of coordination number (required for the bond valence analysis) is usually unambiguous, but that for ternary systems the often continuous distribution of distances observed makes this determination somewhat arbitrary.

It has also been shown that the chemical shift range associated with a particular coordination number and atom can be fairly large (for example, Na coordinated by six O atoms is given by Koller \textit{et al.}\textsuperscript{5} as ranging from –20 to +10 ppm in the materials studied). This is attributed to the dependence of the chemical shift upon the particular chemical environment of the coordinating atoms themselves, \textit{i.e.}, the elements to which they themselves are bonded. It is noticeable that when the surrounding oxygens are bonded to less electronegative elements the chemical shift is in general more positive. Despite the large coordination number of Na in the perovskite materials, the electronegativities of Nb and Ta are reasonably low, perhaps therefore resulting in a more positive chemical shift. This demonstrates how any correlation between chemical shift and structural parameter is dependant on the detail of the number and type of coordinating atom and also on the more longer range structural features. The chemical nature of the solid as a whole strongly affects the chemical shift and it appears that next-nearest neighbour interactions must also be considered in the heavier transition-metal oxides we have studied.

Only a few other materials with high (>8) sodium coordination number have been studied using \(^{23}\text{Na}\) MAS NMR. These include the minerals nepheline (–5.5 and –19.5 ppm for 8 and 9 coordinate sodium, respectively), microcline (–24.3 ppm for 9 coordinate Na) and tourmaline (–13 ppm for 9 coordinate Na) and the material \(Na_3LiYSi_6O_{15}\) (–18 ppm for 9 coordinate Na).\textsuperscript{7} Fluorine containing materials studied include the perovskite \(NaBaLiNiF_6\) (for which a Na chemical shift of <0 ppm may be inferred from the spectra presented,\textsuperscript{41} and the alkali fluorites cryolite, \(Na_3AlF_6\) (2.4 and –9.3 ppm for 6 and 8 coordinate sodium)\textsuperscript{38,42} and chiolite, \(Na_3AlF_6\) (–6 and –21 ppm for 6 and 8 coordinate sodium).\textsuperscript{9} In all cases, the general trend that larger numbers of more distant coordinating atoms results in a decrease in chemical shift was apparent, but that detailed correlations of shift with structural parameters is far from trivial. It should be noted that many other A-site sodium perovskites contain paramagnetic metal...
ions, for example the tungsten bronzes Na₄WO₆, the U(IV) material NaUMo₅, and the fluorides NaMF₃ (M = V, Mn, Co), so measurement of ²³Na MAS NMR spectra is not straightforward.

In conclusion, we have used ²³Na MAS NMR, in conjunction with DFT calculations to aid spectral interpretation, for the study of two perovskite materials, NaTaO₃ and NaNbO₃, where Na is found in an unusually high coordination number environment. The high-resolution achieved by MQMAS was able to determine that our as synthesised sample contained a Pbcm so measurement of P from the two-dimensional spectra were small, in agreement with many literature observations that the chemical shift tends to decrease when Na is coordinated by a larger number of atoms at greater distances. However, a search of the literature provides evidence that detailed correlation with structural information at greater distances is possible. The isotropic chemical shifts we extracted from the two-dimensional spectra were small, in agreement with many literature observations that the chemical shift tends to decrease when Na is coordinated by a larger number of atoms at greater distances. However, a search of the literature provides evidence that detailed correlation with structural parameters is difficult and more long range effects must be taken into account.

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