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Enhanced efficiency of solid-state NMR investigations of energy materials using an external automatic tuning/matching (eATM) robot



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ABSTRACT

We have developed and explored an external automatic tuning/matching (eATM) robot that can be attached to commercial and/or home-built magic angle spinning (MAS) or static nuclear magnetic resonance (NMR) probeheads. Complete synchronization and automation with Bruker and Tecmag spectrometers is ensured via transistor-transistor-logic (TTL) signals. The eATM robot enables an automated "on-the-fly" re-calibration of the radio frequency (rf) carrier frequency, which is beneficial whenever tuning/matching of the resonance circuit is required, e.g. variable temperature (VT) NMR, spin-echo mapping (variable offset cumulative spectroscopy, VOCS) and/or in situ NMR experiments of batteries. This allows a significant increase in efficiency for NMR experiments outside regular working hours (e.g. overnight) and, furthermore, enables measurements of quadrupolar nuclei which would not be possible in reasonable timeframes due to excessively large spectral widths. Additionally, different tuning/matching capacitor (and/or coil) settings for desired frequencies (e.g. ⁷Li and ³¹P at 117 and 122 MHz, respectively, at 7.05 T) can be saved and made directly accessible before automatic tuning/matching, thus enabling automated measurements of multiple nuclei for one sample with no manual adjustment required by the user. We have applied this new eATM approach in static and MAS spin-echo mapping NMR experiments in different magnetic fields on four energy storage materials, namely: (1) paramagnetic ⁷Li and ³¹P MAS NMR (without manual recalibration) of the Li-ion battery cathode material LiFePO₄; (2) paramagnetic ¹⁷O VT-NMR of the solid oxide fuel cell cathode material La₂NiO_{4+ δ_1^2} (3) broadband ⁹³Nb static NMR of the Li-ion battery material BNb₂O₅; and (4) broadband static ¹²⁷I NMR of a potential Li-air battery product LilO₃. In each case, insight into local atomic structure and dynamics arises primarily from the highly broadened (1-25 MHz) NMR lineshapes that the eATM robot is uniquely suited to collect. These new developments in automation of NMR experiments are likely to advance the application of in and ex situ NMR investigations to an ever-increasing range of energy storage materials and systems. © 2016 The Authors. Published by Elsevier Inc. This is an open access article under the CC BY license (http:// creativecommons.org/licenses/by/4.0/).

1. Introduction

The development of solid-state materials for energy storage and conversion (*e.g.* in batteries, supercapacitors, and fuel cells) relies on understanding fundamental relationships between structure and bulk properties such as electronic and ionic conductivity [1-5]. As a direct probe of the environments of atomic nuclei, solid-state nuclear magnetic resonance (NMR) can report on local structure and dynamics [6-8] with relevance to understanding

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electronic and structural phase transitions [9–12] and ionic diffusion [13–16] on a wide range of time scales.

Among the major challenges in acquiring and interpreting solid-state NMR spectra of functional energy materials are the presence of extremely large anisotropic interactions that cannot be effectively averaged by standard techniques such as magic-angle spinning (MAS) or specialized pulse programs [17,18]. Although these interactions provide important electronic and structural details about the system of interest, they also give rise to broad powder patterns that exceed the NMR probe bandwidth (*ca.* 1 MHz), prohibiting the excitation of the entire spectrum in a single experiment. Acquisition of the overall lineshape thus, in the simplest approach, requires measurements at different radio frequency (rf) carrier frequencies, with frequent and time-

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consuming manual readjustment (tuning and matching) of the rf circuit [19]. Extreme broadening effects may arise for several reasons, which we highlight in turn:

- *Paramagnetic interactions.* Systems containing unpaired electrons are subject to hyperfine interactions such as the isotropic Fermi contact shift and the anisotropic electron–nuclear dipolar coupling interaction, leading to large shifts and broadened lineshapes [20]. These shifts are typically on the order of several thousand ppm, although shifted resonances of nearly 10,000 ppm are not unknown [21,22]. Nonetheless, large paramagnetic shifts and anisotropic broadening values provide important structural information, as they can often be rationalized on the basis of spin-transfer (bond) pathways [23].
- Quadrupolar coupling interactions. The coupling of the local electric field gradient (EFG) with the nuclear electric quadrupole moment can lead to exceptionally broad static powder patterns (*ca.* 10 MHz). Particularly problematic quadrupolar nuclei include ^{35/37}Cl, ^{47/49}Ti, ⁵⁹Co, ^{69/71}Ga, ^{79/81}Br, ⁸⁷Sr, ⁹¹Zr, ⁹³Nb, ¹¹⁵In, and ¹²⁷I, with several others (*e.g.* ^{177/179}Hf, ¹⁸¹Ta) considered of limited utility due to extreme broadening [18,24–26]. However, broad quadrupolar lineshapes are highly characteristic and correlate sensitively with local geometry.
- Large NMR shift ranges of heavier spin-½ nuclei. The NMR spectra of heavier spin-½ nuclides span a wide chemical (and also Knight) shift range [27,28]. Chemical shift ranges (and chemical shift anisotropies, CSAs) of 1000–5000 ppm that are exceptionally responsive to the chemical environment have been reported for ¹⁰⁹Ag, ¹¹³Cd, ^{117/119}Sn, ¹²⁵Te, ¹⁸³W, ¹⁹⁵Pt, ¹⁹⁹Hg, and ²⁰⁷Pb [18,28–30].
- Variable temperature (VT-NMR) measurements. The tuning and matching of the probe must be optimized at each temperature in a typical VT-NMR experiment [31–33]. Martin and Zilm [34] ascribe this effect to significant temperature-dependent changes in sample conduction and/or magnetic properties, problems that are also more pronounced for materials that undergo a phase transition over the VT range.
- Presence of multiple NMR-active nuclei. Energy storage and conversion materials often contain several NMR-active isotopes. In these systems, a nuclei-switching technique to record multinuclear spectra could be desirable. Examples of previous (manually re-tuned) multinuclear NMR studies potentially validating this approach include ⁷Li and ³¹P NMR ($v_0 = 117$ MHz and 122 MHz at 7.05 T) of the Li-ion battery cathode materials LiMPO₄ (M = Fe, Mn, Co, Ni) [35,36] and the Li solid electrolyte Li₄SiO₄–Li₃PO₄ [37]; ²³Na, ⁵¹V, and ²⁷Al NMR ($v_0 = 79.3$ MHz, 78.9 MHz, and 78.2 MHz at 7.05 T) of sodium metavanadates [38,39] and glass ceramics [40,41]; and ²⁰⁷Pb and ¹²⁷I NMR ($v_0 = 62.8$ MHz and 60.0 MHz at 7.05 T) of the superionic conductor PbI₂ [42] and of hybrid halide perovskite materials, *e.g.* (CH₃NH₃)PbI₃ [43].

Several specialized techniques have been developed for overcoming these problems and acquiring NMR signals with spectral widths >250 kHz, together referred to as "ultra-wideline NMR spectroscopy" [18]. By far the simplest approach is to sweep the rf carrier frequency, in so-called "frequency-sweep" measurements (also referred to as variable-offset cumulative spectroscopy, VOCS, or spin-echo mapping) [19,44–47]. Alternatively, the use of adiabatic or chirped rf pulses, such as wideline, uniform rate, smooth truncation (WURST) pulses or short high-power adiabatic pulses (SHAP), leads to larger excitation bandwidths at a single carrier frequency, relative to typical high-power rectangular pulses [48–51]. The utility of the adiabatic pulse approach has been extended through sequences such as QCPMG (quadrupolar Carr–Purcell–Mei boom-Gill) echo trains [52-54] and DANTE (Delays Alternating with Nutation for Tailored Excitation) schemes [55–57] to further enhance sensitivity and selectivity in paramagnetic, quadrupolar and large CSA systems. Slow-turning methods, such as the STEA-MER (Slow Turning Echo Amplitude Modulation and Echo Reduction) [58] and STREAQI (Slow Turning Reveals Enormous Anisotropic Quadrupolar Interactions) [59] sequences, also resolve broad quadrupolar parameters at a single carrier frequency, although they are often limited in their practical application by requirements of off-magic angle or right-angle spinning. Finally, various hardware-based solutions are available: solenoid microcoils (and magic-angle coil spinning) to access very high rf field amplitudes [60,61]; the use of weak pulsed field gradients in solution-state NMR for broadband decoupling [62,63]; and magnetic field (rather than rf frequency) sweep experiments, using 0.5–1.0 T superconducting sweep coils [64.65] as well as highfield resistive magnets [66].

In this study, we report on the development and application of an external automatic tuning/matching (eATM) robot to repeatedly and accurately recalibrate the resonance circuit without manual intervention, e.g. during time-consuming experiments requiring collection of many individual sub-spectra. This eATM device has been developed through an ongoing collaboration between the Grey group at the University of Cambridge (UK) and NMR Service GmbH (Erfurt, Germany), and is now commercially available from the latter [67]. The focus of the present work is the initial application of the eATM prototype to enhance the throughput of solidstate NMR studies of various energy storage systems, as tested through our joint effort. We show that use of the eATM robot significantly boosts the overall collection efficiency of broadband, wideline and VT-NMR experiments, allowing overnight acquisition without direct supervision, and with only minor changes to existing protocols and pulse programs. Rather than replacing existing ultra-wideline NMR schemes, the eATM system accommodates and extends these approaches, as we demonstrate with a combined eATM and WURST-QCPMG experiment.

The eATM robot is physically attached to the tuning/matching rods of the NMR probe and can therefore interface with a broad range of probeheads and NMR spectrometers under static as well as MAS conditions. Previously, a similar device optimized for magnetic resonance imaging (MRI) applications reported by Hwang et al. employed a double-acting hydraulic system [68]. Here, our focus is on acquiring broadband (>1 MHz) NMR spectra of solid-state materials, and an electric motor assembly is instead used to drive the rotation of the tuning/matching rods. Koczor et al. [69] have also recently reported an external remotely-controlled tuning/matching device, which still however requires manual readjustment; in the present work, the eATM system dynamically recalibrates the rf circuit prior to each experiment, with no user input, in response to the forward (fwd) and reflected (rfd) power of a low-power continuous wave (cw) rf pulse.

In this contribution, we describe the initial set-up and practical operation of the eATM robot from a user's point of view, its frequency sweep and recalibration capabilities (Section 4.1), report its use in four proof-of-concept studies of various energy storage and conversion materials (Sections 4.2–4.5), and discuss its applications and future developments (Sections 4.6 and 5). The materials have been chosen to highlight the difficulty in acquiring NMR spectra of systems that are paramagnetic, or contain lineshapes significantly influenced by first- and second-order quadrupolar coupling, in particular under static conditions. In an initial section, we first discuss the technological relevance of these materials as well as open questions regarding structure and dynamics that are amenable to solid-state NMR methods (Section 2).

2. Materials systems investigated

2.1. LiFePO₄

The Li-ion battery cathode material LiFePO₄ (LFP) is a wellestablished model system for understanding Li-ion intercalation reactions. In particular, recent studies of nanosized LFP electrodes have established the fundamental role of metastable kinetic intermediates in enabling high-rate battery operation [70–72]. The formation of these intermediates on charging and discharging can in principle be followed with both ⁷Li and ³¹P NMR as two distinct probes of the local structure, providing information complementary to X-ray diffraction and microscopy studies [73]. Moreover, as ⁷Li and ³¹P possess similar Larmor frequencies (117 and 122 MHz, respectively, at 7.05 T), LFP presents an ideal system for testing fully automated nuclei switching. As with other transition-metal phosphates, the ⁷Li and ³¹P NMR shifts are dominated by large paramagnetic interactions-which can be explicitly quantified through bond pathway decomposition analysis [23]and thus high-power adiabatic pulses and/or VOCS-like techniques [19] are typically required to map out the broadband ⁷Li and ³¹P spectra and extract relevant NMR parameters.

2.2. La₂NiO_{4+δ}

Solid oxide fuel cells (SOFCs) are solid-state electrochemical conversion devices that traditionally operate at or above 800 °C. An outstanding goal in current SOFC research is to enable functionality at much lower temperatures through the development of materials that show good oxide-ion conduction below 600 °C. Mixed ionic–electronic conductors (MIECs) in particular help to overcome slow oxygen reduction kinetics in electrode operation at lower temperatures.

The perovskite-derived MIEC La₂NiO_{4+ δ} demonstrates remarkably high oxide-ion conductivity and acceptable electronic conductivity at intermediate temperatures, and is recognized as an important candidate cathode material for next-generation SOFCs [74]. The large oxygen excess (δ) in La₂NiO_{4+ δ} arises from non-stoichiometric incorporation of interstitial oxygen; these interstitial defects are highly mobile and a knock-on interstitialcy mechanism has been proposed to account for the oxide-ion conduction [75].

In our group we have employed La₂NiO_{4+ δ} as a model system to extend ¹⁷O MAS solid-state NMR spectroscopic techniques to paramagnetic oxides. Paramagnetic ¹⁷O NMR in the solid-state is virtually unknown, but in a significant advance Kong et al. have shown the utility of spin-echo mapping in acquiring ¹⁷O spectra for transi tion-metal-containing coordination compounds [21]. A similar broadband approach is required for La₂NiO_{4+ δ}, where ¹⁷O resonances in both the diamagnetic (interstitial sites) and paramagnetic (axial and equatorial sites) shift ranges are observed, spanning a total of *ca.* 1 MHz (at 16.4 T) [76].

Many studies have employed ¹⁷O solid-state NMR experiments at variable temperatures as a probe of thermally-activated local dynamics in oxide-ion conductors, allowing for calculation of exchange rates and relevant kinetic parameters [13,77]. For La₂NNiO_{4+δ}, an orthorhombic-to-tetragonal phase transition at roughly 130 °C has been reported [78], well within the feasible range of VT-NMR at reasonable MAS rates with standard, commercial probes. Indeed, clear changes in lineshapes and intensity of the ¹⁷O VT-NMR spectra of La₂NiO_{4+δ} occur as oxide-ion motional processes become activated near 130 °C [76]. From a technical point of view, however, spin-echo mapping experiments become tedious if performed at many different temperatures; approaches to reduce collection time and improve overall experimental efficiency are welcome.

2.3. B-Nb₂O₅

Niobium oxides [79,80] and Nb-doped compounds [81,82] are emerging in novel energy storage applications, especially for high-power and high-rate applications. While many polymorphs of Nb₂O₅ are known, B-Nb₂O₅ is a crystallographically interesting model oxide compound due to the presence of a single niobium site with distorted octahedral coordination [83,84]. The structure is monoclinic with *C*2/*c* symmetry; it is composed of edgesharing octahedral dimers in a corner-sharing network that is related to the titania polymorphs of rutile and TiO₂(B). While ⁹³Nb NMR would seem an ideal technique to study the local distortions of the octahedral niobium site, the large quadrupolar coupling constants associated with this nucleus necessitate the use of more involved ultra-wideline experiments.

2.4. LiIO₃

Iodine finds use in various energy-related systems, such as the I^-/I_3^- redox couple used in dye-sensitized solar cells [85], lead halide perovskite-based solar cells [43], and as an adduct in Li- O_2 batteries [86]. Recently, Liu et al. have shown that addition of lithium iodide to aprotic Li- O_2 batteries leads to highly reversible cycling and negligible electrode passivation [86]. In this system, iodine functions as a tunable redox mediator, and iodine oxyanions are speculated to play an important role in the electrochemistry. As ¹²⁷I NMR may provide valuable mechanistic insights into this and other systems, we have chosen LiIO₃ as a relevant model compound to study the feasibility and utility of ultra-wideline (static) ¹²⁷I NMR experiments.

3. Experimental section

3.1. Sample preparation

3.1.1. LiFePO₄

Carbon-coated LiFePO₄ was synthesized as previously described [71,73] according to the solid-state reaction of Kobayashi et al. [87]. In detail, 0.556 g lithium carbonate (Li₂CO₃, Sigma–Aldrich, 99.997%), 2.681 g iron(II) oxalate dihydrate (Fe(II)C₂O₄·2H₂O, Sigma–Aldrich, 99%), 1.714 g ammonium dihydrogen phosphate (NH₄H₂PO₄, Sigma–Aldrich, 99.999%), and 0.261 g Ketjen black (EC-600JD AkzoNobel) were high energy ball-milled for 40 min to obtain homogeneously mixed precursors. The precursors were then pressed into a pellet and sintered at 600 °C for 6 h under an atmosphere of flowing Ar.

3.1.2. La₂NiO_{4+δ}

Samples of La₂NiO_{4+ δ} were prepared via the solid-state reaction route described previously [88–90]. Stoichiometric amounts of La₂O₃ (Alfa Aesar, REacton, 99.999%; pre-dried) and NiO (Aldrich, 99.999%) were mixed in a mortar and pestle, isostatically pressed, sintered in air at 1300 °C for 6–12 h, and ground. Several intermediate sintering and grinding steps were repeated until phase purity was attained, as determined by laboratory powder X-ray diffraction. Samples of ¹⁷O-enriched La₂NiO_{4+ δ} were prepared by heating the as-synthesized powder (0.1–0.3 g) at 1000 °C under an atmosphere of 70% ¹⁷O₂ (Cambridge Isotope Labs, used as received) in a sealed quartz tube for 24 h. Samples were slowly cooled (1 °C min⁻¹) from 1000 °C to ensure maximal uptake of ¹⁷O.

3.1.3. B-Nb₂O₅

 $B-Nb_2O_5$ was prepared *via* thermal oxidation of NbO₂ (Alfa Aesar, 99.5+%) at 850 °C for 24 h in air. Rietveld refinement of the

powder X-ray diffraction data [79] showed a minor (*ca.* 2%) contamination from the high-temperature H-Nb₂O₅ polymorph.

3.1.4. LiIO₃

LiIO₃ (97%) was obtained from Sigma-Aldrich. Powder X-ray diffraction did not show any impurities (Supporting Information, Fig. S2) and the sample was used as received.

3.2. CASTEP calculations and simulations of NMR spectra

3.2.1. B-Nb₂O₅

First principles, periodic density functional theory (DFT) calculations of the NMR parameters of B-Nb₂O₅ were performed in the plane-wave code CASTEP (v. 8.0) with ultrasoft pseudopotentials generated 'on-the-fly' [91–94]. The magnetic shielding tensor was calculated within the gauge-including projector-augmented wave (GIPAW) approach while the quadrupolar parameters were calculated directly from the principal components of the electric field gradient at the nucleus. Electron exchange and correlation were treated with the Perdew-Burke-Ernzerhof (PBE) functional within the generalized gradient approximation (GGA) [95]. Calculations were performed with a plane-wave kinetic energy cutoff of 600 eV and k-point sampling of the Brillouin zone with a Monkhorst–Pack grid [96] finer than $2\pi \times 0.06 \text{ Å}^{-1}$. Convergence of isotropic shielding and quadrupolar coupling was checked with respect to both basis-set energy and k-point sampling. Extraction of the Euler angles between the chemical shielding and electric field gradient tensors was performed using MagresView [97]. LiNbO₃, with an experimental isotropic shift of -1004 ppm [98], was used to determine the reference shielding for ⁹³Nb. Euler angle and tensor conventions are given in Supporting Information, Table S1.

3.2.2. LiIO3

A similar computational methodology was adopted as for B-Nb₂O₅, with calculations performed using a plane-wave kinetic energy cutoff of 700 eV and k-point sampling with a $8 \times 8 \times 8$ Monkhorst–Pack grid [96]. Ultrasoft pseudopotentials were generated on-the-fly using the zeroth-order regular approximation (ZORA) as implemented in CASTEP [99]. Euler angle and tensor conventions are given in Supporting Information, Table S2. ¹²⁷I chemical shielding tensors were referenced to the ¹²⁷I chemical shift of Nal (226.71 ppm).

3.3. Solid-state NMR experiments

All NMR experiments were performed with the eATM robot attached to the tuning/matching rods of various Bruker probes, and the eATM controller interfaced with Bruker Avance III consoles. The pulse sequences were modified by insertion of a transistor-transistor-logic (TTL) preamble to trigger the synchronization of the auto tuning/matching algorithm (Fig. S1), using a low-power cw pulse of 0.01 W, prior to the start of each data collection. Bruker TopSpin 2.1 and 3.2 were used for raw data processing.

3.3.1. ⁷Li and ³¹P MAS NMR of LiFePO₄

Solid-state ⁷Li and ³¹P MAS NMR experiments were performed on a 7.05 T Bruker Avance III 300 MHz spectrometer ($v_0 = 116.6$ MHz and 121.5 MHz for ⁷Li and ³¹P, respectively) using a Bruker 2.5 mm HFX MAS probe at a spinning speed of 30 kHz. Spin-echo mapping spectra were recorded at room temperature using a rotor-synchronized Hahn echo pulse sequence of the form ($\pi/2$)_x- τ -(π)_y- τ -acquire with a pulse length of 4.17 µs ($\pi/2$ pulse, optimized using LiF and NH₄H₂PO₄ secondary references) at an inherent rf field strength of *ca*. 60 kHz, and recycle delays of 0.25 s and 50 ms for ⁷Li and ³¹P, respectively. For each nucleus, a total of 11 sub-spectra were recorded in steps of 500 ppm (58.3 kHz for ⁷Li and 60.7 kHz for ³¹P), from 116.35 to 116.93 MHz for ⁷Li and from 121.62 to 122.22 MHz for ³¹P. The ⁷Li and ³¹P chemical shifts were referenced to LiF at –1 ppm and NH₄H₂PO₄ at 1 ppm, respectively [100,101].

3.3.2. ¹⁷O MAS VT-NMR of $La_2NiO_{4+\delta}$

Solid-state ¹⁷O MAS NMR experiments were carried out on a 16.4 T Bruker Avance III 700 MHz spectrometer ($v_0 = 94.99$ MHz for ¹⁷O) using a Bruker 4 mm HXY MAS probe. Spin-echo mapping experiments (step size of 1000 ppm, *i.e.* 95 kHz, with 12 or 13 recorded sub-spectra) were performed at a MAS frequency of 12.5 kHz using a rotor-synchronized Hahn echo pulse sequence of the form ($\pi/6$)_x- τ -($\pi/3$)_y- τ -acquire, with a pulse length of 2.43 µs ($\pi/6$ for liquid H₂O) at an inherent rf field strength of *ca.* 34 kHz, and a recycle delay of 20 ms. Temperature calibration was performed in a separate MAS experiment using the known temperature dependence of the ²⁰⁷Pb resonance of Pb(NO₃)₂, with an accuracy of ±5 °C [102,103]. The ¹⁷O VT-NMR spectra were collected on ¹⁷O-enriched samples packed in ZrO₂ rotors with ZrO₂ caps. ¹⁷O chemical shifts were externally referenced to H₂O at 0.0 ppm at room temperature.

3.3.3. ⁹³Nb static NMR of B-Nb₂O₅

Solid-state ⁹³Nb NMR experiments were collected on a 16.4 T Bruker Avance III 700 MHz spectrometer ($v_0 = 171.3$ MHz for ⁹³Nb) using a Bruker 4 mm HXY probe. Static Hahn-echo mapping experiments were recorded at room temperature from 167.8 to 174.8 MHz in 41 steps of *ca.* 1000 ppm (170 kHz). Given the effect of strong quadrupolar coupling on nutation, the rf pulse was directly optimized on the B-Nb₂O₅ sample, yielding an effective $\pi/2$ pulse length of 1.0 µs. For each of the 41 spectra, 1664 scans, temporally spaced by a 1 s recycle delay, were co-added and Fourier transformed. LiNbO₃, with an isotropic shift of –1004 ppm, was used as a secondary shift reference [98].

3.3.4. ¹²⁷I static NMR of LiIO₃

Solid-state ¹²⁷I NMR experiments were carried out on a 16.4 T Bruker Avance III 700 MHz spectrometer ($v_0 = 140.1$ MHz for ¹²⁷I) using a Bruker 4 mm HXY MAS probe. Frequency-stepped static spectra were acquired at room temperature using the WURST-QCPMG broadband excitation scheme [51] employing an 8-step phase cycle and a WURST-80 pulse length of 50 µs [49]. For each slice, two spectra were taken with opposing WURST sweep directions to compensate for relaxation during the sweep. An experimentally optimized rf field strength of 10.4 kHz was used, with a frequency step of 0.75 MHz. Spikelet spacing in the QCPMG pulse was 8196 Hz with an echo delay of 50 µs and a recycle delay of 0.5 s. The ¹²⁷I spectra were referenced to NaI powder at 227 ppm [104].

4. Results and discussion

4.1. The eATM system

The eATM system comprises a robot that is placed underneath the probehead (already inside the bore of the magnet) and subsequently connected to an eATM controller (Fig. 1a and b). In addition to the eATM robot itself (Fig. 1c), the equipment includes specially designed rods with adjustable clamping connectors ("fingertips") that can be attached to the outside of the tuning/matching rods of any commercial or homebuilt static or MAS probehead (Fig. 1d). These height-adjustable rods connect to piezoelectric engines inside the cover of the robot (Fig. 1c). After connecting



Fig. 1. (a) Schematic showing the interplay between the eATM system and NMR spectrometer enabling static or MAS NMR experiments with measurement-synchronized automatic tuning and/or matching. (b) Photos of the eATM robot installed underneath 4.0 mm MAS probeheads at a 200 MHz (4.7 T) and a 700 MHz (16.4 T) magnet. (c) Close-up photograph of the eATM robot during operation. (d) Close-up of the connection between the length-adjustable brass rods and the probehead made *via* clamping connectors ("fingertips"), with various diameters for different types of tuning/matching rods.

the eATM rods to the probe, the robot is itself secured to a plastic floor plate of the appropriate length to prevent self-rotation (Fig. 1b). In this way, the system can fully interface with various probe designs as well as magnets with differences in antivibration leg setups and floor plate thicknesses.

The piezoelectric engines are furthermore connected to actuators that are powered by the eATM controller, which makes use of the fwd and rfd power of a low power cw tuning/matching pulse for adjusting the rf carrier to a specific NMR frequency prior to the actual NMR experiment (Fig. 1a). The fwd and rfd power is shunted via a bidirectional coupler, which is incorporated into the rf circuit between the NMR probehead and spectrometer. The eATM control software determines the standing wave ratio (SWR) based on fwd and rfd power at the desired NMR carrier frequency. An automatic ("on-the-fly") recalibration of the rf circuit is then performed algorithmically through robot-controlled changes of the tuning and/or matching capacitors/coils of the probehead, with simultaneous recalculation of the SWR. The loop is repeated until a SWR minimum is reached, corresponding to optimal tuning/matching of the rf circuit to the carrier frequency. The complete automatic tuning/matching procedure (from initial trigger to start of the NMR acquisition) typically requires less than 20 s. Full synchronization of the eATM system and the NMR spectrometer is enabled with TTL input and output signals for start and stop, respectively, via a preamble in the applied pulse sequence that defines this synchronization (Supporting Information, Fig. S1a). This preamble for automatic tuning/matching can be readily implemented into any pulse sequence for Bruker and Tecmag consoles and is possible for others.

In the following, we will discuss four applications of the eATM robot in static and MAS spin-echo mapping experiments of different kinds of energy storage and conversion materials.

4.2. ⁷Li and ³¹P MAS NMR of LiFePO₄

Using the eATM system, a single fully automated measurement of ⁷Li and ³¹P MAS NMR of LiFePO₄ has been performed (Fig. 2), comprising 11 sub-spectra in carrier frequency steps of 500 ppm for each nucleus. Given approximately 40 min per sub-spectrum, the entire measurement requires *ca*. 15 h overnight, with no user adjustment required.

As seen in Fig. 2c, this approach allows automatic collection of signal over a broadband range of *ca.* 10 MHz. In practice, the first sub-spectrum for each nucleus was collected by initially recalling the nucleus-specific, pre-saved tuning/matching positions corresponding to the appropriate base carrier frequency (as described in Section 4.6), and then further optimizing the rf circuit by running the eATM autotune algorithm. Subsequent sub-spectra were collected using a typical frequency sweep/VOCS method, as necessitated by the highly broadened (2000–3000 ppm) spinning sideband manifolds for both ⁷Li and ³¹P. The paramagnetic lineshapes are in excellent agreement with previous reports [23,105,106], showing that the eATM system enables more efficient acquisition of spectra of several nuclei concurrently, without loss of broadband spectral fidelity, allowing for accurate extraction of hyperfine shifts and dipolar coupling parameters.

4.3. ¹⁷O high-temperature MAS NMR of $La_2NiO_{4+\delta}$

Spin-echo mapped ¹⁷O NMR spectra of La₂NiO_{4+ô} at two different sample temperatures, below (79 °C) and above (148 °C) the reported orthorhombic-to-tetragonal phase transition, have been measured (Fig. 3). While the temperatures have been set manually, the collection of NMR spectra is automated via the eATM device. Due to the limited excitation bandwidth of the Hahn echo experiments, each broadband spectrum requires the acquisition and summation of 12–13 sub-spectra with rf carrier frequency step size of 1000 ppm (95 kHz) to reproduce the features of all diamagnetic and paramagnetic oxygen environments accurately. Temperature-dependent changes in the resonance circuit necessitate a probe re-tuning step after heating to 148 °C, which has also been automatically carried out by the eATM robot.

The main changes seen at high temperature (148 °C) are (1) the disappearance of the diamagnetic, lower frequency resonance (535 ppm) assigned to interstitial oxide defects and (2) a narrowing of the resonance at 5400 ppm assigned to equatorial oxygen sites within the perovskite layers. The former is associated with the onset of interstitial motion that leads to a large decrease in the signal intensity; the latter change is attributed to a less distorted (more symmetrical) equatorial environment with a smaller hyperfine shift dispersion, as expected in the high-temperature tetragonal phase [76,107]. Further work is in progress to correlate the low-temperature onset of interstitial oxide-ion dynamics in La₂NiO_{4+ δ} with local structural displacements and distortions of the other (paramagnetic) oxygen environments.

4.4. ⁹³Nb static NMR of B-Nb₂O₅

Despite some favorable properties such as 100% natural abundance and relatively high gyromagnetic ratio, solid-state ⁹³Nb



Fig. 2. (a) ⁷Li MAS NMR spectrum of LiFePO₄ at 7.05 T with a MAS speed of 30 kHz (black trace, offset), acquired with spin-echo mapping by summing 11 sub-spectra (colored traces) with rf carrier frequency step size of 500 ppm (58.3 kHz). (b) ³¹P MAS NMR spectrum of LiFePO₄ under same conditions (black trace, offset), acquired by summing 11 sub-spectra (colored traces) with rf carrier frequency step size of 500 ppm (60.7 kHz). (c) Overall broadband ⁷Li and ³¹P MAS NMR spectrum of LiFePO₄ spanning > 5 MHz, acquired in a single fully automated overnight measurement using the eATM system. (Sub-spectra are depicted as colored traces as in (a) and (b).)



Fig. 3. ¹⁷O MAS VT-NMR spectra of ¹⁷O-enriched La₂NiO_{4+ $\delta_{1}}$, acquired using spin-echo mapping by summing 12–13 sub-spectra (colored traces) with rf carrier frequency step size of 1000 ppm (95 kHz). Spectra were collected at sample temperatures below (79 °C) and above (148 °C) the orthorhombic–tetragonal phase transition of La₂NiO_{4+ $\delta_{1}}$ (roughly 130 °C). Spectra were obtained at 16.4 T with a MAS rate ω_{r} of 12.5 kHz. Asterisks denote spinning sidebands.</sub></sub>

NMR studies are limited by the large nuclear quadrupolar coupling constants of distorted niobium environments, which can exceed 100 MHz. Although obtaining acceptable signal-to-noise levels for a single ⁹³Nb spectrum requires only *ca*. 30 min, manually adjusting the matching and tuning over the course of one day in order to capture the entire quadrupolar lineshape is inefficient and often impractical. Using the eATM device, we obtained the broadband ⁹³Nb spectrum of B-Nb₂O₅ (Fig. 4), spanning a range of 6.97 MHz, by summing 41 individual Hahn-echo sub-spectra automatically acquired in 20 h. Fitting the features and discontinuities of the broadband B-Nb₂O₅ ⁹³Nb NMR spectrum, aided by *ab initio* calculations, enabled the extraction of accurate magnetic shielding and quadrupolar tensors (Supporting Information, Table S1). The

fit was performed within the Solid Lineshape Analysis (SOLA) program in TopSpin. Initial parameter values were taken from DFT calculations based on the reported X-ray structure [84]. Given the relatively small CSA, in comparison to the quadrupolar coupling, the relative orientation between these two tensors was fixed at the DFT derived value, while the other parameters were varied to achieve the best fit. Estimated errors are proportional to the sensitivity of the fit to each parameter. These quadrupole tensor values ($C_Q = 46(1)$ MHz, $\eta_Q = 0.62(3)$; *cf.* DFT values of -42.25 MHz/0.60 and -39.1 MHz/0.47 for the fixed and relaxed structures, respectively) reflect the bonding and significant local distortions present in the NbO₆ octahedra of this second-order Jahn–Teller distorted d⁰ niobium(V) oxide.



Fig. 4. ⁹³Nb static NMR spectrum of B-Nb₂O₅ at 16.4 T (v_0 = 171.3 MHz). The overall spectrum (black trace) is the summation of 41 individual sub-spectra (colored traces). DFT calculations of NMR tensors and Euler angles were performed to guide the fit (red trace, see main text), using quadrupole coupling parameters of C_Q = 46 (1) MHz and η_Q = 0.62(3) (Section 3.2.1; Supporting Information, Table S1). The sub-spectra have been acquired via spin-echo mapping with an rf carrier frequency step size of *ca*. 1000 ppm (170 kHz); the overall spectrum spans *ca*. 7 MHz.

4.5. ¹²⁷I static NMR of LiIO₃

Despite the high natural abundance (100%) and relatively large gyromagnetic ratio (γ = 5.39 × 10⁷ rad s⁻¹ T⁻¹) of the ¹²⁷I nucleus, both solution and solid-state ¹²⁷I NMR studies are rare on account of the extremely strong quadrupolar nature $(I = 5/2, 0 = -71 \text{ fm}^2)$ of this nuclide [108]. Most previous solid-state NMR studies have been performed on systems of simple metal iodides such as NaI, PbI₂, or alkaline earth metal iodides [42,109]. As systems such as iodates (IO_3) are expected to exhibit large quadrupole coupling constants due to the lower symmetry about the iodine nucleus, we have tested the applicability of the eATM in measuring very broad, multiple frequency-stepped ¹²⁷I spectra. Fig. 5 shows the summed ¹²⁷I solid-state NMR spectrum of LiIO₃ using the frequency-stepped WURST-QCPMG pulse sequence. Multiple frequency-stepped slices spanning nearly 25 MHz can be automatically acquired by the eATM robot without manual adjustment. To compare to DFT results, the lineshape was simulated with the QUEST software [110] to correctly account for the large quadrupolar interaction. Simulation using DFT-calculated parameters of $C_0 = 970.1 \text{ MHz}$ and $\eta_0 = 0.0$ (Supporting Information, Table S2) qualitatively agrees with the ultra-wideline experiment, showing an extremely broad second-order quadrupolar lineshape. In particular, the experimental spectrum exhibits discontinuities at 176 and 153 MHz, in excellent agreement with the relevant subsection of the simulated broadband spectrum. A detailed ¹²⁷I NMR study reporting a complete fitting of the experimental spectrum is in progress [111]. Clearly, this investigation shows the utility of eATM experiments in capturing and assigning lineshapes spanning tens of MHz with broadband excitation schemes.

4.6. Discussion

Application of the eATM robot is advantageous whenever multiple (automatic) recalibrations of the rf circuit are required, *e.g.* frequency-stepped [44,45,57] (VOCS [14,21,46,112] and/or spinecho mapping [113–116]) experiments, VT measurements, and/or *in situ* NMR experiments, especially in cases where changes in sam-



Fig. 5. ¹²⁷I static NMR spectrum of LiIO₃, acquired using WURST-QCPMG. The experimental spectrum (black) is a sum of 64 individual sub-spectra (32 slices). The inset shows a simulation of the NMR spectrum using DFT-calculated NMR parameters of C_Q = 970.1 MHz and η_Q = 0.0 (Section 3.2.2; Supporting Information, Table S2).

ple properties can influence the optimal rf setting [73]. Besides straightforward "on-the-fly" automatic tuning/matching to a desired carrier frequency, the eATM system also permits saving different carrier frequencies as tuning/matching servo positions. Upon sending a position-specific TTL signal to the controller *via* the pulse sequence preamble (Supporting Information, Fig. S1b–e), the eATM servos are rotated to the saved values, which is automatically followed by a further recalibration step to the exact rf carrier. This functionality permits fully automated switching between nuclei, provided the pre-set carrier frequencies are within the frequency range of the probe hardware, *e.g.* ⁷Li and ³¹P, where $B_0 = 7.05$ T, at 116 and 122 MHz, respectively (Section 4.2).

The flexibility of the system is demonstrated by its integration with a variety of probe and magnet designs, spectrometer systems, and in principle all pulse programs. Moreover, successful operation is observed even for older probes with significant tuning/matching hysteresis of the SWR. The efficiency of the eATM approach is ensured by the speed of the recalibration step, which allows for rapid collection of numerous sub-spectra. This is of particular relevance in the field of energy storage materials, where many ex situ samples (e.g. as a function of lithiation) may be studied in a single experiment; for measurements spanning a wide spectral range, without the eATM system the overnight magnet time would often be wasted on a single carrier frequency. Finally, the eATM device enables a range of experiments normally considered impractical, such as spanning the entirety of extremely broadened quadrupolar lineshapes, or searching for transitions in nuclear quadrupole resonance (NQR) spectroscopy where the resonance often lies at an unknown position within a *ca.* 10 MHz range.

5. Conclusions

We have designed an external automated tuning/matching (eATM) robot system and report on its applications to ultrawideline and broadband solid-state NMR experiments for a wide variety of probes, consoles, pulse sequences and materials applications. The robot allows for rapid tuning/matching with a broad (>25 MHz) tuning/matching range, permits more efficient frequency-sweep experiments in off-peak hours (especially overnight), and enables automatic multinuclear and VT NMR studies not possible with other ultra-wideline techniques. Ongoing research focuses on the implementation of other adiabatic pulse sequences analogous to those previously used in ultra-wideline quadrupolar NMR [18] and broadband paramagnetic NMR [57], as well as the combined use of pulsed field gradients (PFG) in performing automated multinuclear diffusion studies. The alternative use of the eATM robot for off-MAS or satellite transition magic angle spinning (STMAS) NMR (by accurately adjusting the magic angle rod) is in progress, as well as the development of automated NQR or hybrid NMR-NQR studies [117]. Furthermore, the combined application of (1) automated sample re-orientation in static NMR experiments using a goniometer with (2) re-calibration of the rf circuit via the eATM is likely to become a powerful approach to study (A)BMS effects, which is particularly of interest for energy storage materials [6,118]. Finally we note the potential of "upgrading" static in situ NMR probeheads with eATM, given the importance of compensating for changes in battery materials during cycling that influence the optimal tuning/matching settings [73,119]. We conclude that this contribution and future developments in eATM devices are highly promising for extending in situ and ex situ solid-state NMR studies of energy materials.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.jmr.2016.12.008. Data supporting this publication are available at www.repository.cam.ac.uk.

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