Inorganic Chemistry

An Investigation of Chlorine Ligands in Transition-Metal Complexes via 35Cl Solid-State NMR and Density Functional Theory Calculations

Christopher A. O'Keefe,[†] Karen E. Johnston,[†] Kiplangat Sutter,[‡] Jochen Autschbach,[‡] Régis Gauvin,[§] Julien Trebosc, ́ § Laurent Delevoye,§ Nicolas Popoff, [∥] Mostafa Taoufik,[∥] Konstantin Oudatchin,[⊥] and Robert W. Schurko*,[†]

† Department of Chemistry an[d](#page-15-0) Biochemistry, University of Windsor, Windsor, Ontario N9B 3P4, Canada

‡ Department of Chemistry, University at Buffalo, State University of New York, Buffalo, New York 14260, United States

§ UniversitéLille Nord de France, CNRS UMR8181, Unitéde Catalyse et de Chimie du Solide, UCCS USTL, F-59655 Villeneuve d'Ascq, France

_∥
Laboratoire de Chimie, Catalyse, Polymères et Procédés (UMR-C2P2-5265 CNRS/ESCPE-Lyon/UCBL) ESCPE Lyon, F-308-43, Boulevard du 11 Novembre 1918, F-69616 Villeurbanne Cedex, France

[⊥]Steacie Institute for Molecular Sciences, National Research Council, 100 Sussex Drive, Ottawa, Ontario K1A OR6, Canada

S Supporting Information

[AB](#page-15-0)STRACT: [Chlorine liga](#page-15-0)nds in a variety of diamagnetic transitionmetal (TM) complexes in common structural motifs were studied using ³⁵Cl solid-state NMR (SSNMR), and insight into the origin of the observed 35Cl NMR parameters was gained through first-principles density functional theory (DFT) calculations. The WURST-CPMG pulse sequence and the variable-offset cumulative spectrum (VOCS) methods

were used to acquire static ³⁵Cl SSNMR powder patterns at both standard (9.4 T) and ultrahigh (21.1 T) magnetic field strengths, with the latter affording higher signal-to-noise ratios (S/N) and reduced experimental times (i.e., <1 h). Analytical simulations were performed to extract the $35Cl$ electric field gradient (EFG) tensor and chemical shift (CS) tensor parameters. It was found that the chlorine ligands in various bonding environments (i.e., bridging, terminal-axial, and terminal-equatorial) have drastically different ³⁵Cl EFG tensor parameters, suggesting that ³⁵Cl SSNMR is ideal for characterizing chlorine ligands in TM complexes. A detailed localized molecular orbital (LMO) analysis was completed for NbCl₅. It was found that the contributions of individual molecular orbitals must be considered to fully explain the observed EFG parameters, thereby negating simple arguments based on comparison of bond lengths and angles. Finally, we discuss the application of ³⁵Cl SSNMR for the structural characterization of WCl₆ that has been grafted onto a silica support material. The resulting tungsten-chloride surface species is shown to be structurally distinct from the parent compound.

ENTRODUCTION

Group V and VI transition metals (TM) are commonly used to fabricate a wide range of complexes with various uses, such as reagents in redox reactions, $1-4$ antitumor agents, $5-7$ and superconducting materials.^{8−10} However, perhaps their most ubiquitous use is for the cataly[si](#page-15-0)s [o](#page-15-0)f reactions involvin[g](#page-15-0) [or](#page-15-0)ganic and organometallic species.[11](#page-15-0)−[16](#page-15-0) Their multiple oxidation states and relatively large atomic sizes allow for the coordination of a variety of ligands and the s[yn](#page-15-0)t[he](#page-15-0)sis of catalysts that can be finetuned for specific purposes. Such complexes find use in both homogeneous and heterogeneous catalytic processes; the latter is widely thought to afford higher selectivity and efficiency.^{17,18} As a result, the design and application of heterogeneous catalysts, which involves the chemical or physical immob[iliza](#page-15-0)tion of the TM species on high-surface area materials, represent burgeoning areas of research.^{19−21}

Structural characterization of heterogeneous catalysts is often difficult, as the support mate[ria](#page-16-0)l[s a](#page-16-0)re almost always disordered and insoluble, thereby limiting or preventing their analysis by conventional techniques such as X-ray diffraction (XRD) or solution-state NMR spectroscopy. Surface techniques such as X-ray photoelectron spectroscopy $(XPS)^{22}$ and IR spectrosco $py^{23,24}$ have previously been used to study immobilized species; however, the structural information is oft[en](#page-16-0) limited. Solid-state N[MR](#page-16-0) (SSNMR) spectroscopy is now widely used for molecular-level characterization of heterogeneous catalysts,^{25−28} providing information on structural motifs, bonding, interatomic distances, symmetry and ground-state electronic distr[ibutio](#page-16-0)ns. Furthermore, unlike XRD methods, SSNMR is capable of probing short-range order in systems lacking longrange order (i.e., amorphous and/or disordered solids).

SSNMR experiments involving ${}^{1}H$ and ${}^{13}C$ are the most common for studies of heterogeneous catalysts.27,29−³¹ However, our research group has shown that it may be possible to investigate such systems from the perspecti[ve of](#page-16-0) t[he](#page-16-0)

Received: April 29, 2014 Published: August 27, 2014

Table 1. 35Cl NMR Parameters Obtained from Spectra Acquired at 21.1 T

sample	site	$ C_0 ^a$ (MHz)	$\eta_{\text{Q}}^{\text{b}}$	$\delta_{\rm iso}{}^c$ (ppm)	Ω^d (ppm)	κ^e	α^f (deg)	β (deg)	γ (deg)
α -WCl ₆		21.00(10)	0.029(5)	800(40)	200(100)	1.0(2)	0(10)	0(10)	0(10)
β -WCl ₆		21.00(10)	0.029(5)	800(40)	200(100)	1.0(2)	0(10)	0(10)	0(10)
WOCl ₄		26.05(10)	0.115(5)	620(50)	550(150)	$-1.0(4)$	50(5)	15(10)	50(5)
MoOCl ₄		30.50(10)	0.585(10)	650(50)	800(100)	1.0(2)	0(5)	90(5)	0(10)
	$\mathfrak{2}$	25.30(10)	0.510(5)	950(100)	800(150)	1.0(2)	0(5)	90(5)	90(5)
	3	25.00(10)	0.500(5)	800(50)	300(100)	1.0(3)	0(10)	90(10)	0(10)
	$\overline{4}$	23.30(5)	0.540(20)	730(30)	235(50)	0.0(5)	0(20)	0(10)	0(20)
NbCl ₅		25.30(5)	0.385(10)	250(150)	460(300)	1.0(2)	90(30)	0(10)	90(30)
	2	15.00(8)	0.081(7)	850(50)	200(200)	0.0(5)	0(20)	0(5)	0(30)
	3	14.70(9)	0.655(5)	990(40)	300(200)	0.0(3)	45(15)	0(20)	45(15)
TaCl _s		25.70(8)	0.430(10)	260(100)	250(200)	1.0(5)	90(30)	0(10)	90(30)
	2	15.65(15)	0.043(6)	640(50)	300(100)	1.0(3)	0(30)	0(5)	0(30)
	3	16.70(30)	0.480(20)	900(40)	700(50)	$-1.0(3)$	0(30)	90(5)	0(30)
WCl_6-SiO_2		25.98(5)	0.11(1)	450(30)	300(100)	$-1.0(5)$	50(10)	15(5)	50(10)
a_{C} α α β β	b_{11} (\mathbf{r})	T^2 λ / T^2 c _S	$\sqrt{2}$ $\sqrt{2}$ $\sqrt{2}$	$\ln d_{\Omega}$ \sim	$c \quad e_{11} \quad \gamma/c$	$S \cap \Omega$	f_{TL} , f_{L}		

 ${}^aC_Q = eQV_{33}/h$. ${}^b\eta_Q = (V_{11} - V_{22})/V_{33}$. ${}^c\delta_{iso} = (\delta_{11} + \delta_{22} + \delta_{33})/3$. ${}^d\Omega = \delta_{11} - \delta_{33}$. ${}^e\kappa = 3(\delta_{22} - \delta_{iso})/\Omega$. *f* The Euler angles, α , β , and γ , define the relative orientations of the EFG and CS tensors. The "ZYZ" convention for rotation is used herein, as described by Dye et al.,⁸³ and as implemented in the WSolids⁵⁸ software package.

S[che](#page-16-0)me 1. Schematic Representation of (A) Tungsten(VI) Chloride (WCl₆), (B) Tetrachlorotungsten(VI) [Ox](#page-16-0)ide (WOCl₄) and Tetrachloromolybdenum(VI) Oxide (MoOCl₄), (C) Niobium(V) Chloride (NbCl₅) and Tantalum(V) Chloride (TaCl₅), and (D) the Different Chlorine Bonding Environments in the Bridging Pentahalide Species^a

^aBridging chlorine atoms (Cl_{br}) are shown in blue, terminal-axial (Cl_{ax}) in green and terminal-equatorial (Cl_{eq}) in orange.

TM nuclei; to date, we have characterized bulk metallocene complexes utilizing $91Zr$, $47/49Ti$, $93Nb$, and $139La SSNMR$.^{32–36} The spectra of many of these nuclei are very broad, ranging from several hundred kHz to MHz in breadth; according[ly,](#page-16-0) [S/](#page-16-0) N tends to be very poor in many of these spectra, which suggests limitations for studying the diluted, supported TM species in heterogeneous catalysts. However, recent developments in broadband excitation sequences and other signal enhancing techniques, $37-39$ along with the increasing availability of higher magnetic field strengths and improved NMR probes and hardware, s[ugges](#page-16-0)t that SSNMR of such systems may hold much promise. Unfortunately, for certain TM nuclei such as 95/97Mo, 103Rh, 181Ta, and 183W, probing the metal center using SSNMR currently remains challenging due to their inherently unreceptive natures. In such cases, it is often more convenient and informative to conduct NMR experiments

involving other nuclei on the ligands surrounding the metal center, since their NMR parameters are likely to be sensitive to structural differences. A prime example of this is chlorine, which is found widely in many TM complexes.

Chlorine has two naturally occurring NMR-active isotopes, 35Cl and 37Cl, with natural abundances of 75.78 and 24.22%, respectively. Both have nuclear spins of $I = 3/2$ and low gyromagnetic ratios ("low-γ"), with relatively small quadrupole moments of -8.165 and -6.435 fm^{2,40} ³⁵Cl is the preferred . target NMR nuclide due to its higher receptivity (i.e., both its natural abundance and Larmor freque[ncy](#page-16-0) are higher).

The $35Cl$ quadrupolar interaction (QI), which arises from coupling between the ³⁵Cl nuclear quadrupole moment and local electric field gradients (EFGs) at the ³⁵Cl nucleus, is very sensitive to subtle differences in structure. Accordingly, the quadrupolar coupling constant, C_{Q} , and the asymmetry parameter, $\eta_{\rm Q}$ (see Table 1 for definitions), which are determined from simulations of the ³⁵Cl SSNMR powder patterns, reflect variance in str[uc](#page-1-0)ture and bonding. ³⁵Cl SSNMR spectra typically exhibit central transition $(CT, +1/2 \leftrightarrow -1/2)$ powder patterns that are strongly influenced by the secondorder quadrupolar interaction (SOQI); the two satellite transition patterns (ST, $\pm 3/2 \leftrightarrow \pm 1/2$) are predominantly broadened by the first-order quadrupolar interaction (FOQI) and are rarely observed in ³⁵Cl SSNMR spectra (excepting in cases where the magnitude of C_Q is very small).

The majority of ³⁵Cl SSNMR studies to date have focused on systems with the chlorine atoms at the center of ground-state electronic distributions of high spherical or Platonic symmetry (e.g., Cl[−] ions weakly coordinated by water molecules and other ligands, $ClO₄⁻$, etc.). Such environments are generally associated with small values of C_Q and have relatively narrow $35C1$ CT powder patterns that can be acquired using conventional Hahn-echo experiments and high-power rectangular pulses.41−⁴⁴ Examples of such systems include metalchloride salts, hydrochloride (HCl) salts of amino acids, and pharmaceuti[cal](#page-16-0)s[. T](#page-16-0)he application of ³⁵Cl SSNMR to systems in which Cl is involved in covalent bonding is very rare by comparison, since the values of C_Q are much larger and the powder patterns much broader. Some recent works include investigations of group 13 chloride salts, 45 chlorine-containing metallocenes,⁴⁶ organometallic transition metal complexes,⁴⁶ and organic compounds with covalent [C](#page-16-0)−Cl bonds.⁴⁸ All of these studie[s](#page-16-0) report the use of high magnetic fields a[nd](#page-16-0) specialized pulse sequences due to the exceptionally b[roa](#page-16-0)d ³⁵Cl powder patterns associated with these systems. Such spectra, and the concomitant experiments necessary for their acquisition, fall under the classification of ultrawideline (UW) NMR spectroscopy.^{49,50}

Herein, we describe the application of $35C1$ SSNMR in conjunction with fi[rst-](#page-16-0)principles density functional theory (DFT) calculations to study diamagnetic inorganic chlorinecontaining TM complexes (Scheme 1) that exhibit a variety of common structural motifs and are representative of several broader classes of TM complexes wi[de](#page-1-0)ly used in catalysis. All of these compounds have been previously characterized by 35° Cl NQR spectroscopy; this allows us to determine whether the values of C_O extracted from simple analytical simulations under the high-field approximation (i.e., $\nu_0 > \nu_Q$) are valid, and to also accurately measure the values of η_{Q} , which are unavailable from NQR spectra of spin-3/2 nuclides. First, we demonstrate the application of the wideband uniform-rate smooth truncationquadrupolar Carr−Purcell−Meiboom−Gill (WURST-CPMG)

pulse sequence^{38,51} for the rapid acquisition (i.e., $<$ 1 h) of UW ³⁵Cl SSNMR spectra at both 9.4 and 21.1 T. In one case, the use of ³⁵Cl N[MR an](#page-16-0)d ³⁵Cl NQR experiments in tandem for the differentiation of polymorphs is explored. Second, quadrupolar parameters determined from simulations of these spectra are compared to those obtained from plane-wave DFT calculations. This data, along with theoretically derived ³⁵Cl EFG tensor orientations, are used to propose relationships between the ³⁵Cl quadrupolar parameters and the nature of the chlorine structural environments (i.e., the identity of the metal (M), M−Cl bond lengths, geometric arrangements, etc.). Third, a full analysis of the contributions of the individual molecular orbitals to the principal components of the 35Cl EFG tensor for $NbCl₅$ is presented and utilized to explore the origins of the experimentally observed ³⁵Cl quadrupolar parameters. Fourth, we present an example of an application of ³⁵Cl SSNMR to a tungsten chloride species grafted onto a silica support material. On the basis of experimental and theoretical data on the bulk species presented herein, it is possible to make a rough structural interpretation of this result. Finally, we discuss the potential applications of this methodology for the study of heterogeneous TM chloride catalysts.

EXPERIMENTAL AND COMPUTATION DETAILS

Sample Preparation. Compounds α -WCl₆, NbCl₅, and TaCl₅ were purchased from Strem Chemicals, and compounds $WOCl₄$, and MoOCl4 were purchased from Sigma-Aldrich Canada, Ltd. All samples were used without further purification or modification (phase purity was checked with powder XRD, vide infra). β -WCl₆ was prepared by the slow sublimation of the α -form under a nitrogen environment.⁵ All of the compounds were stored and packed under an inert nitrogen atmosphere in a dry glovebox due to their air- and moisture-sensiti[ve](#page-16-0) natures. All samples were finely ground and packed into shortened 5 mm outer diameter glass NMR tubes and then capped and sealed with Parafilm to limit their exposure to atmosphere for the duration of the experiments.

Preparation of WCl₆-SiO₂. A toluene solution of WCl₆ was added to a toluene suspension of dehydroxylated silica (Degussa Aerosil 250, pretreated at 700 °C under 10[−]⁵ bar vacuum), and the resulting mixture was stirred overnight at room temperature. After filtration of the supernatant, the solid was washed three times with toluene, one time with pentane and dried under a vacuum (10[−]⁵ bar). The amount of tungsten in the sample is 5.48 wt %. HCl evolution was checked by IR spectroscopy as in a previous communication.⁵³

Powder X-ray Diffraction (PXRD). PXRD experiments were performed at 100 K on a Bruker Apex 2 Kappa [Di](#page-16-0)ffractometer using graphite monochromatized Mo K_a radiation (λ = 0.7107 Å). Theoretical PXRD patterns were simulated for all samples using the Powder Cell software package.⁵⁴

³⁵Cl SSNMR Spectroscopy. A Varian Infinity Plus spectrometer equipped with a wide-bore Ox[for](#page-16-0)d 9.4 T magnet $(\nu_0(^1H) = 400 \text{ MHz})$ and ν_0 (³⁵Cl) = 39.16 MHz) was used for the acquisition of static (i.e., stationary samples) 35Cl SSNMR spectra for all samples. The experiments were conducted on a Varian Chemagnetics 5 mm triple-resonance (HXY) T3 MAS NMR probe. Spectra were also collected using a Bruker Avance II spectrometer with a 21.1 T $(\nu_0(^{35}Cl) = 88.2 \text{ MHz})$ standard-bore magnet at the National Ultrahigh-field NMR Facility for Solids in Ottawa, Ontario, Canada. A home-built 4 mm double resonance probe was used for all experiments. The spectrum of WCl_6-SiO_2 was acquired on a Bruker Avance III spectrometer with an 18.8 T standard-bore magnet using a 3.2 mm HX MAS probe at UniversitéLille Nord de France. Chemical shifts were referenced to 1 M NaCl (aq), with NaCl(s) ($\delta_{\rm iso}$ = 0 ppm) used as a secondary reference.

All ³⁵Cl CT powder patterns were too broad to be uniformly excited with a single, rectangular, high-power pulse. Hence, the WURST-CPMG pulse sequence^{38,51} was used with a 50 μ s WURST pulse swept over 2000 kHz (sweep rate of 40 MHz/ms) for experiments at 9.4 and 21.1 T, whereas at 18.8 T the WURST pulse was swept over 4000 kHz (sweep rate of 80 MHz/ms). The experiments at 18.8 T also used a continuous-wave ¹H decoupling field of 30 kHz that was applied for the duration of the scan, including during the WURST pulses. A detailed list of the experimental parameters can be found in the Supporting Information (Tables S1−S3). Spectra were acquired using the variable offset cumulative spectrum (VOCS) technique⁵⁵⁻⁵⁷ with each subspectrum acquired using a transmitter offset of 100 kHz. [Subspectra](#page-15-0) [were](#page-15-0) [proces](#page-15-0)sed using the NUTS program fr[om A](#page-16-0)corn software. The free induction decays (FIDs) of the subspectra were processed with digital filtering, Fourier transformation, and a magnitude calculation. The individual subspectra were then coadded (9.4 T) or skyline projected (21.1 T) to produce the total spectrum. The spectra consist of a series of narrow lines, called spikelets, whose outer manifold is representative of the shape of the powder pattern. Analytical simulations of the spectra were generated using WSolids⁵⁸ (9.4 T) and QUEST⁵⁹ (21.1 T) software packages; precise values of the quadrupolar parameters are obtained by matching the positions [of](#page-16-0) the key spectral disc[ont](#page-16-0)inuities (i.e., "shoulders" and "horns") and not the positions of the individual spikelets. Uncertainties in the quadrupolar parameters were estimated by bidirectional variation of each parameter and visual comparison of experimental and simulated spectra.
³⁵Cl transverse relaxation time constants (T_2) were measured using

the CPMG pulse sequence. Echo trains were acquired by placing the transmitter at each of the spectral discontinuities. The intensity of each echo train was then plotted as a function of time and fit to an

³⁵Cl NQR Spectroscopy. Experiments were conducted on a Varian Chemagnetics triple resonance (HXY) T3 MAS probe with a 5 mm coil and a Varian Infinity Plus console. The probe was placed as far away from the magnet as possible to limit the effects from the external magnetic field. The Hahn-echo pulse sequence was used with a 3.5 μ s 90° pulse width and a recycle delay of 0.5 s. Processing of the ³⁵Cl NQR spectra was done using NUTS.

Plane-Wave DFT Calculations. ³⁵Cl EFG and CS tensor parameters were calculated using the $CASTEP⁶⁰$ plane-wave density functional theory code in the Materials Studio 5.0 software package. Calculations employed the generalized gra[di](#page-16-0)ent approximation (GGA), the gauge-including projector augmented wave (GIPAW) formalism and revised Perdew, Burke and Ernzerhof (rPBE) functionals, with the core-valence interactions being described by on-the-fly pseudopotentials (Table S4, Supporting Information). Integrals over the Brillouin zone were performed using a Monkhorst-Pack grid with a k-point spacing of 0.07 Å[−]¹ . Wave functions were expanded in pla[ne](#page-15-0) [waves](#page-15-0) [with](#page-15-0) [a](#page-15-0) [kinetic](#page-15-0) [energy](#page-15-0) [cuto](#page-15-0)ff of 300 eV. Crystal structures were obtained from the Inorganic Crystal Structure Database (ICSD), and calculations were completed both prior to and post geometry optimization of the structures (i.e., all of the atomic positions were allowed to vary simultaneously, while the cell parameters were fixed). All calculations were performed on the Shared Hierarchical Academic Research Computing Network (SHARCNET) using either 8 or 16 cores and 16 GB of memory

³⁵Cl Natural Bond Orbital/Natural Local Molecular Orbital **Analysis of NbCl₅.** All computations were performed using a developer's version of Amsterdam Density Functional (ADF)⁶¹ package. Experimental single-crystal XRD crystal structures were used as starting structures for geometry optimization. For t[he](#page-16-0) computation of EFG tensor parameters and the geometry optimization routines, relativistic effects were included by utilizing the zeroth-order regular approximation (ZORA) in its scalar form.⁶² All calculations were carried out using the revised Perdew−Burke−Ernzerhof (rPBE) functional.63,64 The triple-ζ doubly polarized ([TZ](#page-16-0)2P) all-electron Slater-type basis sets were used for Nb and Cl in all computations. Localized [mol](#page-16-0)ecular orbital (LMO) analyses of ³⁵Cl EFGs were performed as described previously.⁶⁵ LMOs were generated using the natural bond orbital (NBO) algorithm⁶⁶ implemented in a locally modified version of the NBO 5.0 program.⁶⁷ The set of "natural" LMOs generated by the NBO code was used for the EFG analysis.

■ RESULTS AND DISCUSSION

For ease of discussion, all complexes have been grouped according to their structural similarities (Scheme 1). Some of the complexes exist as polymorphs in the solid state, which are pure substances that crystallize in more than one [st](#page-1-0)able phase, with each having a distinct arrangement of atoms.⁶⁸ Powder Xray diffraction (pXRD) patterns, which are useful for determining the presence of known crystalline p[ha](#page-16-0)ses, as well as for detecting for the presence of multiple polymorphs and crystalline impurities, were acquired for all of the samples (see Supporting Information, Figures S7−S11).

Before each group of systems is discussed in detail, it is useful [to comment on some of the common fea](#page-15-0)tures observed for all of the 35Cl SSNMR spectra, as well as some of the aspects of the plane-wave DFT calculations of the 35Cl EFG tensor parameters. First, all ³⁵Cl SSNMR spectra were acquired at both standard (9.4 T) and ultrahigh magnetic field strengths (21.1 T). The higher field strength serves to increase S/N due to (i) a more favorable Boltzmann distribution and (ii) the reduction in the breadth of the SOQI-dominated CT pattern by an amount proportional to the inverse of B_0 . Despite the general superiority of the 21.1 T data in terms of increased S/ N, reduced experimental times and improved resolution, we note that the experiments completed at 9.4 T were crucial for parametrizing the experiments and optimizing our use of the 21.1 T spectrometer. The spectra acquired at 21.1 T are shown in the manuscript, while those acquired at 9.4 T are presented in the Supporting Information. The only exception to this was β -WCl₆, which was only studied at 9.4 T.

Sec[ond, the breadth of eac](#page-15-0)h powder pattern is extremely large (700−1750 kHz) due to broadening of the CT by the SOQI. For all of the systems discussed herein, the SOQI has a much larger effect on the CT powder pattern than the chlorine chemical shift anisotropy (CSA); hence, the spectral features arise almost solely from the former. As such, the values obtained for the isotropic chemical shift ($\delta_{\rm iso}$), span (Ω), skew (κ), and the Euler angles $(\alpha, \beta, \text{ and } \gamma)$ describing the relative orientations of the EFG and CS tensors (see Table 1 for definitions) have large uncertainties associated with them. The chlorine CS tensor parameters were calculated, and the r[es](#page-1-0)ults are shown in the Supporting Information, along with a figure demonstrating that the inclusion of CSA has little effect on the appearance of the [powder patterns \(Table](#page-15-0) S7, Figure S17).

Third, the extreme breadths of the CT powder patterns render it difficult to distinguish chlor[ine environments tha](#page-15-0)t are chemically similar (i.e., those with comparable bond lengths and angles) but crystallographically distinct. However, the specific types of bonding environment for chlorine (i.e., bridging, terminal-axial, or terminal-equatorial) are easily discernible from the ³⁵Cl SSNMR spectra. Furthermore, techniques such as ³⁵Cl NQR and T_2 editing of CPMG echo trains generally allow for the resolution of chemically similar environments and the accurate measurement of the NMR parameters.

Fourth, higher intensity than expected is always observed in the higher frequency range (i.e., leftmost regions) of each spectrum in comparison to the low frequency (rightmost) range. This effect is more pronounced for certain samples in comparison to others probably due to effects such as T_2 anisotropy and/or variation in probe/circuit characteristics Table 2. Experimental and Calculated (Using CASTEP)³⁵Cl EFG Tensor Parameters and Comparison of Predicted NQR Frequencies Obtained from ³⁵Cl SSNMR Data to Those Reported in the Literature^a

 a Calculation of the NMR parameters was completed both prior to and after the geometry optimization of the structure. b The NQR frequencies were calculated from NMR parameters obtained from solid-state ³⁵Cl SSNMR spectra using $\nu_Q^{NQR} = (C_Q/2)\sqrt{(1 + (\eta_Q)^2/3)}$.

over the broad range of acquisition frequencies (vide infra). Fifth, in some cases, it is possible to acquire signal extending past the edges of the CT in each powder pattern. This signal is attributed to the ³⁵Cl satellite transition (ST) patterns (i.e., $-3/$ $2 \leftrightarrow -1/2$ and $+1/2 \leftrightarrow +3/2$, which are visible owing to the high abundance of chlorine in each complex. ST patterns are broadened immensely by the first-order quadrupolar interaction over a range of several MHz and often are associated with T_2 ⁽³⁵Cl) parameters that are distinct from those of the CT. In the case of 35Cl, the ST patterns result in minimal interference with the CT patterns. We note that the acquisition of the full ST powder pattern is generally unnecessary for UW SSNMR spectra, since the CT pattern provides information on all of the relevant NMR parameters. However, with quadrupolar parameters obtained from simulations of the CT patterns, it is possible to rapidly locate the key discontinuities in the STs and to potentially further refine the quadrupolar parameters.⁵⁷

Despite the extreme breadths of the CT powder patterns, it was possible to attain high S/N, high-resolution spectra [in](#page-16-0) relatively short time frames (e.g., 32 s to 26 min at 21.1 T) owing to the broadband excitation of the WURST pulses, the long ³⁵Cl transverse relaxation times constants (T_2) that favor CPMG-type experiments, and the high chlorine concentrations in these samples.

As mentioned, chlorine sites that are chemically very similar are not easily differentiated in the ³⁵Cl SSNMR spectra, and

therefore analytical simulations are representative of the convolution of the quadrupolar parameters of these sites. By fitting the simulations to the positions of the well-defined discontinuities of the spectra, it was possible to extract the $35³⁵$ Cl EFG tensor parameters C_Q and η_Q (Table 1). ³⁵Cl NQR measurements have previously been conducted on all of the samples discussed herein. Since NQR measurem[en](#page-1-0)ts on spin-3/ 2 nuclei involve the measurement of a single transition, no values of η_{Q} are typically reported, which is a limitation of the NQR experiment.^{69,70} Nonetheless, we see generally good agreement between our experimental values of C_Q and those from NQR measu[reme](#page-16-0)nts on ³⁵Cl nuclei possessing values of $\eta_{\rm O}$ near to zero (i.e., axially symmetric EFG tensors, Table 2).

The quadrupolar parameters determined from ³⁵Cl NMR spectra, in conjunction with ³⁵Cl EFG tensor parameters obtained from DFT calculations, were used to differentiate chlorine sites in distinct bonding arrangements (i.e., bridging (Cl_{br}) , terminal-axial (Cl_{ax}) , and terminal-equatorial (Cl_{eq}) , vide infra). No consideration is given to the theoretical chlorine nuclear magnetic shielding tensors herein, since in most cases, the experimentally determined values of the chemical shift tensor parameters are associated with large uncertainties (as discussed above). The 35 Cl EFG tensor parameters were calculated using CASTEP⁶⁰ both prior to and after the geometry optimization of each structure (Table 2). Geometry optimization of the structur[es](#page-16-0) resulted in small changes in bond lengths (Table S5, Supporting Information), large changes in the computed EFG tensor parameters and greatly reduced forces b[etween atoms \(Table S6, Supportin](#page-15-0)g Information). In all cases, it was found that terminal chlorines have negative values of C_{Q} , whereas [bridging chlorines have positive va](#page-15-0)lues. The theoretical magnitudes of C_Q are found to be consistently less than the experimentally measured values (Figure 1), similar to the observations of Rossini et al.⁴⁶ We note that the sign of C_O cannot be measured directly from the ³⁵Cl NMR spectra but is readily determined by the calcul[atio](#page-16-0)ns.

Figure 1. Comparison between the experimental and calculated values of (A) C_Q and (B) η_Q for all sites within the complexes studied. All calculated values were obtained from 35Cl EFG tensor calculations completed on geometry optimized structures using CASTEP. The values of both C_Q and η_Q for the pseudo-bridging chlorine sites in MoOCl4 were omitted due to poor correlation between experiment and calculation.

Tungsten(VI) Chloride (WCl₆). Tungsten(VI) chloride exists as one of two polymorphs, α or β . The α -form has one tungsten atom and one chlorine atom within the asymmetric unit (i.e., there is one unique chlorine environment). There are six chlorine atoms arranged around a central tungsten atom in an octahedral arrangement (Scheme 1). The β -form has two tungsten atoms and three chlorine atoms within the asymmetric unit. The three chlorine sites have si[mi](#page-1-0)lar W−Cl bond lengths (2.23, 2.26, and 2.34 Å) and comparable Cl−W−Cl bond angles (91.3, 91.6, and 90.3°), resulting in a slightly distorted octahedral arrangement of chlorine atoms about the central tungsten atom.⁷¹ Both forms feature hexagonal close packed lattices of chlorine atoms with tungsten ions filling the octahedral hol[es.](#page-16-0) The β -form can be prepared from the α form by the slow sublimation of the α -form under $N_2(g)$, or annealing a sample at a temperature greater than 200 $^{\circ}$ C (we chose the former procedure, see Experimental and Computational Details for details).⁵²

The ³⁵Cl static SSNMR spectra of α -WCl₆ and β -WCl₆ [acquired at 21](#page-2-0).1 and 9.4 [T,](#page-16-0) respec[tively,](#page-2-0) [are](#page-2-0) [shown](#page-2-0) [in](#page-2-0) [Figure](#page-2-0) [2,](#page-2-0) panels A and B. The latter compound was synthesized after experiments on all other samples had been conducted at 21[.1](#page-6-0) T; for reasons described below, no high-field data for this compound is presented. The CT of the α -form has a breadth of ca. 700 kHz and is characteristic of a powder pattern broadened by the second-order quadrupolar interaction. The discontinuities typical of such a spectrum are clearly visible and the overall manifold of the spikelet pattern is representative of a single distinct chlorine site. This spectrum was best simulated with one site, $C_Q = 21.0 \text{ MHz}$ and $\eta_Q = 0.029 \text{ (Table 1)}.$ This result is consistent with the α -form of tungsten(VI) chloride, which has one distinct chlorine environment. The CT [of](#page-1-0) the β form is ca. 1.5 MHz in breadth (due to the inverse dependence of the CT breadth on the applied external magnetic field strength). Despite the presence of multiple, magnetically and crystallographically distinct chlorine sites in the β -form, there appears to be only a single CT pattern, almost identical to the pattern acquired for α -WCl₆ at 9.4 T (Figure S12, Supporting Information). In fact, this pattern can be subjected to a similar one-site simulation, yielding identical [quadrupolar parameters](#page-15-0) [to those of](#page-15-0) α -WCl₆ (within the limits of their uncertainties).

The benefit of ultrawideline SSNMR experiments for rapidly acquiring spectra of anisotropically broadened powder patterns is tempered by their ability to resolve patterns arising from sites with very similar quadrupolar parameters. Even our attempts to acquire higher-resolution Hahn-echo spectra at key discontinuities failed to aid in clearly resolving the three unique chlorine sites in the β -form (Figure 2C). In this case, ³⁵Cl NQR is clearly very useful for trying to resolve the structurally similar sites in the β -for[m](#page-6-0), as well as confirming the identity of each of these polymorphs. Both the α - and β -forms have previously been studied by ³⁵Cl NQR spectroscopy. The ³⁵Cl NQR spectrum of the α form has a single peak corresponding to a frequency at $\nu_{\rm Q}^{\rm NQR} = 10.520 \text{ MHz}$, whereas that of the β -form has three peaks $(\nu_{\text{Q}_{\text{VOR}}}^{\text{NQR}} = 10.520, 10.525, \text{ and } 10.576 \text{ MHz})$.⁷² The values of $\nu_{\mathbb{Q}}^{\text{NQR}}$ predicted from the experimentally determined ³⁵Cl EFG tensor parameters obtained for WCl₆ (Table 2), using the expression $\nu_{Q}^{NQR} = (C_{Q}/2)(1 + (\eta_{Q}^{2})$ $(3))^{1/2}$, (3) are in good agreement with the previously reported NQR f[req](#page-4-0)uencies. Using the same NMR probe, we conducted our o[wn](#page-16-0) 35Cl NQR experiments, and found a single peak at $\nu_{\rm Q}^{\rm NQR}$ = 10.5154 MHz for α -WCl₆ and three peaks at $\nu_{\rm Q}^{\rm NQR}$ = 10.512, 10.519, and 10.548 MHz for β -WCl₆, in excellent agreement with previous results (Figure 2D,E). The quadrupolar parameters extracted for $β$ -WCl₆ from NQR experiments are so similar, that not even experim[en](#page-6-0)ts at 21.1 T could effectively separate these patterns; hence, no further NMR experimentation was carried out.

It is clear that NMR and NQR are ideal partners for studying polymorphs, especially in cases where there is no a priori knowledge of the quadrupolar parameters. The specificity of the NQR experiment, while valuable for differentiating sites with similar quadrupolar parameters, is also a crutch, in that it results in tedious searches for NQR resonances over broad frequency ranges. However, the UW SSNMR experiments are capable of providing rapid and accurate determinations of quadrupolar parameters, allowing for the NQR experiment to be conducted in a frequency region of high specificity, and forgoing the

Figure 2. Static ³⁵Cl SSNMR spectrum acquired using frequency-stepped WURST-CPMG for (A) α -WCl₆ at 21.1 T and (B) β -WCl₆ at 9.4 T with corresponding analytical simulations shown in red. (C) Hahn-echo experiments conducted on β -WCl₆ at (i) the high and (ii) the low frequency discontinuities. ³⁵Cl NQR spectra for (D) α -WCl₆ and (E) β -WCl₆. The NQR frequencies ($\nu_{\rm Q}^{\rm NQR}$) for each of the sites are shown in the figure. The spectra were acquired with a transmitter frequency of 10.52 MHz.

monotonous, long "search periods" associated with conventional NQR experimentation.

In both α -WCl₆ and β -WCl₆, the C_Q values are at the lower end of the range of values reported for a chlorine directly bound to a transition metal.⁷⁴ As well, the values of $\eta_{\rm O}$ are close to zero, indicating that the ³⁵Cl EFG tensors are nearly axially symmetric in each case a[nd](#page-16-0) that the chlorine atoms are in environments of high cylindrical symmetry. This is expected for chlorine atoms that are in terminal bonding arrangements with respect to the central metal atom and indicates that V_{33} is likely to be oriented directly along or close to the direction of the metal-chlorine bond (vide infra).

Plane-wave DFT calculations of the ³⁵Cl EFG tensors for both the α - and β -forms of WCl₆ are presented in Table 2. In both forms of the compound, the values of C_Q are found to be negative (consistent with the terminal chlorine site) [a](#page-4-0)nd

Figure 3. Theoretical ³⁵Cl EFG tensor orientations in the molecular frames for (A) α -WCl₆ and (B) β -WCl₆ (C) WOCl₄, (D) MoOCl₄ with (i) the pseudo-bridging chlorine sites (Cl1), (ii) the terminal chloride sites opposite the bridging sites (Cl2), and (iii) the terminal chlorine sites adjacent to the bridging chlorine sites (Cl3 and Cl4), and (E) NbCl₅ with (i) $Cl_{b\nu}$ (ii) Cl_{eq} and (iii) Cl_{ac} . All pictured orientations were determined from ³⁵Cl EFG tensor calculations completed on geometry-optimized models using CASTEP.

consistently lower in magnitude than the experimental values. The values of $\eta_{\rm Q}$ are predicted to be near zero, also in agreement with experiment. The calculated ³⁵Cl NMR parameters obtained prior to optimization of the structure are in poorer agreement with experiment, highlighting the need for geometry optimization of the structures. For example, in the set of calculations on model systems of β -WCl₆ without geometry optimization, there is a large range of C_Q values predicted, which is not expected for chlorine environments that are chemically very similar. This suggests that the reported crystal structure of β -WCl₆ may be inaccurate, as geometry optimization of the structure yields quadrupolar parameters that are more consistent with the ³⁵Cl SSNMR data and not subject to wild variations from site to site. It is also important to

note that the predicted quadrupolar parameters are not ideal (i.e., quite far from experimental values) for use as a priori starting points for the acquisition of NQR spectra; however, NQR experiments featuring the use of WURST pulses may be useful in this respect.⁷⁵

The theoretically determined ³⁵Cl EFG tensor orientations for α - and β -WCl₆ a[re](#page-16-0) shown in Figure 3A,B, respectively. In both cases, the largest principal components of each ³⁵Cl EFG tensor, V_{33} , are oriented along, or nearly along the W−Cl bonds $(\angle(V_{33}-Cl-W) = 178^{\circ}$ for α -WCl₆ and 180° for β -WCl₆). The value of $\eta_{\rm O}$ is close to zero in both forms, so the orientations of V_{11} and V_{22} are of little consequence, since $V_{11} \approx V_{22}$.

Tetrachlorotungsten(VI) Oxide (WOCl₄) and Tetrachloromolybdenum(VI) Oxide (MoOCl₄). The crystal

Figure 4. Static ³⁵Cl SSNMR spectra acquired at 21.1 T for (A) WOCl₄ and (B) MoOCl₄. Analytical simulations representative of the entire powder patterns are shown in red, while individual sites are shown in blue (Cl1), green (Cl2), black (Cl3), and orange (Cl4).

structure of $WOCl₄$ has one molecule in the asymmetric unit with four chlorine atoms in magnetically equivalent environments $(r(W–Cl) = 2.285 \text{ Å})^{76}$ The molecule has a square pyramidal structure with an oxygen atom double bonded to the central tungsten atom at the [ape](#page-16-0)x (Scheme 1). The chlorine atoms sit in a square plane, which lies slightly below the plane of the tung[s](#page-1-0)ten atoms. Previous XRD studies of $WOCl₄$ have suggested that the molecules are associated through weak interactions between the oxygen of one molecule and the tungsten of another. The molecules stack on top of one another to form linear arrays with asymmetric W−O−W bridges (the intramolecular W−O distance is 1.737 Å, whereas the intermolecular W−O distance is 2.258 Å).

The local molecular structure of $MoOCl₄$ is similar to that of $WOCl₄$ (Scheme 1); however, their extended structures are strikingly different. Neutron and X-ray diffraction studies have shown that $MoOCl₄$ contains two molecules in the asymmetric unit with four chemically and magnetically distinct chlorine

sites. The square pyramidal units do not form linear arrays, but instead, are associated through weak interactions between the chlorine atoms of one molecule and the molybdenum atom of another (the Mo−Cl bond length is ca. 2.3 Å, and the intermolecular Mo \cdots Cl distance is 2.9 Å). These intermolecular interactions cause the molecules to form quasi-dimeric structures, with a single chlorine from each molecule forming the *pseudo-bridge* between adjacent pyramidal units.⁷

The ³⁵Cl SSNMR spectra for WOCl₄ and MoOCl₄ are shown in Figure 4, panels A and B, respectively. The [C](#page-16-0)T of the WOCl4 pattern has a breadth of ca. 1075 kHz. The appearance of the discontinuities indicates that there is only one chlorine environment within the complex, in agreement with the aforementioned structure. The markedly different appearance of the 35 Cl SSNMR spectrum for MoOCl₄ clearly reflects the structural differences between the two complexes. The discontinuities in the $MoOCl₄$ spectrum are less well-defined than those in the $WOCl₄$ spectrum, and it appears that there

Figure 5. Static ³⁵Cl SSNMR spectra acquired at 21.1 T for (A) NbCl₅ and (B) TaCl₅. Corresponding analytical simulations representative of the entire powder patterns are shown in red and simulations of the individual sites are shown in blue (Cl_{br}) , green (Cl_{ax}) , and orange (Cl_{ca}) .

are at least four discernible patterns, three of which are very similar and overlapping with each other. The spectrum for $MoOCl₄$ has a much larger breadth (ca. 1750 kHz) than that of WOCl4 and exhibits notable distortions, with greater intensity apparent in the high-frequency region of the spectrum.

The WOCl₄ spectrum is best simulated with a single site C_O = 26.05 MHz and η_{Q} = 0.115 (Table 1). The MoOCl₄ spectrum is simulated with four overlapping patterns of equal intensity arising from chlorine environm[en](#page-1-0)ts with distinct quadrupolar parameters (Table 1). Three of the patterns have values of C_Q and η_Q that lie within experimental error of one another, indicating that there [a](#page-1-0)re three chlorine sites with similar chemical environments. The resulting overlap of these three patterns causes broadening of the outer discontinuities.

The magnitudes of C_Q for WOCl₄ and MoOCl₄ are considerably larger than that of WCl_6 (Table 1); it is not trivial to explain this difference, given the distinct modes of bonding in and different symmetries of these [t](#page-1-0)wo sets of compounds. In particular, one must be cautious in looking for relationships between M−Cl bond lengths and magnitudes of C_Q . The natures of the M–Cl bonds in WCl₆ are likely to be very different from those in the MOCl₄ species, despite the very similar M−Cl bond lengths (i.e., by natures, we mean the differences in bonds in terms of their covalent/ionic character, hybridization of s and p orbitals, and multiple bond character).⁷⁸ Even the comparison of the structurally similar

WOCl4 and MoOCl4 species does not reveal any simple explanation for the variation in values of C_Q (clearly the nature of the extended structure in the latter case plays a role in influencing the EFG tensor). The value of $\eta_{\rm O}$ for WOCl₄ is comparable to that of WCl_6 and is consistent with previous observations for terminal chlorine environments. The values of $\eta_{\rm O}$ for MoOCl₄ are all moderate, indicating nonaxial EFG tensors and reflecting the association of the molecules into dimeric structures. Indeed, a careful analysis of the contributions from individual molecular orbitals to the EFG tensor is necessary to account for differences in values of C_O , as well as for examining the relationships between intra- and intermolecular contacts and EFG tensor orientations and parameters (vide infra). 65

Both WOCl₄ and MoOCl₄ systems have previously been studied by 35° 35° Cl NQR spectroscopy, and similar results regarding the types and multiplicities of chlorine environments were found. The NQR spectrum for the $WOCl₄$ complex has a single peak corresponding to the single chlorine chemical environment, and the value of the NQR frequency is in good agreement with the experimentally determined value of C_O (Table 2). The spectrum of the $MoOCl₄$ complex has four NQR frequencies, three of which are very similar. The highest frequen[cy](#page-4-0) resonance (ca. 3 MHz higher than the others) was assigned to the chlorine atoms which act as the pseudo-bridge

between the metal centers.⁷² This is consistent with our findings discussed above.

Plane-wave DFT calculati[on](#page-16-0)s were performed for $WOCl₄$ and $MoOCl₄$ using structures obtained from the ICSD. The DFT calculations of the 35 Cl EFG tensors for WOCl₄ predict one magnetically distinct chlorine environment, as expected. The value of C_Q predicted from post-geometry optimized calculations is found to be negative, and $\eta_{\rm Q}$ close to zero, consistent with the chlorine atoms being in a terminal environment. The theoretical magnitudes of C_Q are considerably lower than the experimentally determined values, but the values of η_{Q} are more closely reproduced. ³⁵Cl DFT calculations completed for MoOCl₄ predict four chlorine environments, all with negative values of C_Q and moderate values of η_{O} , consistent with our assignments above.

The ³⁵Cl EFG tensor orientations for WOCl₄ and MoOCl₄ are shown in Figure 3, panels C and D, respectively. Since the value of $\eta_{\rm O}$ is near zero for the single chlorine site in WOCl₄, this indicates that V_{33} V_{33} V_{33} is the distinct component in the ³⁵Cl EFG tensor and that the values of V_{11} and V_{22} are similar in magnitude. Because of the symmetry of the $WOCl₄$ molecule, the presence of the chlorine atoms in a plane below the tungsten atom, and the covalently bound O atom, V_{33} is not oriented exactly along the W−Cl bond (i.e., \angle (V₃₃−Cl–W = 4.45 $^{\circ}$). In MoOCl₄, the three magnetically distinct chlorine sites (the *pseudo-bridging chlorine atom (Cl1)*, the terminal chlorine atom opposite to the pseudo-bridge (Cl2), and the terminal chlorine atom adjacent to the pseudo-bridge (Cl3 and $Cl(4)$) have distinct ³⁵Cl EFG tensor orientations. The designation pseudo-bridging refers to a chlorine atom that has a covalent bond with one metal center, and a distant contact with a second metal center from a neighboring molecule. All three sites have moderate values of η_{Q_l} indicating that their ³⁵Cl EFG tensors are not axially symmetric, which is indicative of an absence of cylindrical symmetry of the ground-state electron density about these chlorine sites. For Cl1, V_{33} is oriented near the Mo−Cl bond (∠(V_{33} –Cl–Mo) = 1.85°) and the value of C_{O} is negative, which is consistent with similar observations for terminal chlorine environments.⁴⁵ It has been shown that chlorine atoms which are involved in true bridging interactions between transition metal cent[ers](#page-16-0) have ³⁵Cl EFG tensor orientations in which V_{33} is oriented perpendicular (or approximately perpendicular) to the plane of the M−Cl−M bonding arrangement and positive values of C_0 ⁴⁶ Sites Cl2, Cl3, and Cl4 all have the V_{33} components directed near the direction of their respective Mo[−](#page-16-0)Cl bonds (\angle (V₃₃−Cl−Mo) = 5.67° and 12.95°, respectively), as is typical of chlorines in terminal bonding arrangements.

Niobium(V) Chloride (NbCl₅) and Tantalum(V) Chloride (TaCl₅). The crystal structure of niobium pentachloride $(NbCl_s)$ has one molecule in the asymmetric unit with nine crystallographically distinct chlorine atoms. The complex forms a dimeric structure in which the chlorine atoms form two octahedra sharing a common edge, with the niobium atoms occupying the centers of the octahedra. This structure results in three specific bonding arrangements for the chlorine atoms: bridging (Cl_{br}) , terminal-axial (Cl_{ax}) , and terminal-equatorial (Cleq) (Scheme 1D). The average Nb−Cl bond distances are 2.555, 2.302, and 2.250 Å for Cl_{br} , Cl_{ax} and Cl_{eq} , respectively.⁷⁹ Tantalum(V) c[hlo](#page-1-0)ride (TaCl₅) is isostructural to NbCl₅ and has average Ta−Cl bond lengths of 2.547, 2.307, and 2.225 [Å](#page-16-0) for Cl_{br} , Cl_{av} and Cl_{eq} , respectively.⁸⁰

The static 35 Cl SSNMR spectra of NbCl₅ and TaCl₅ are shown in Figure 5, panels A and B, respectively. The CT patterns have breadths of ca. 1300 kHz ($NbCl_s$) and ca. 1400 kHz (TaCl₅). Fo[r](#page-9-0) each system, the outer manifolds of the spikelet patterns reveal several clearly defined discontinuities that indicate the presence of three distinct patterns, consistent with the presence of the Cl_{br} , Cl_{ax} , and Cl_{eq} bonding environments (vide infra).

The analytical simulation for the entire powder pattern of $NbCl₅$ is shown in Figure 5A as the red trace, with the individual patterns shown as the blue (C_Q = 25.30 MHz and η_Q = 0.385), green (C_Q = 15.0 [MH](#page-9-0)z and η_Q = 0.081), and orange $(C_Q = 14.70 \text{ MHz}$ and $\eta_{Q} = 0.655)$ traces. We note that the patterns are simulated with 1:1:1 signal intensity (instead of the expected 1:2:2, $Cl_{\text{br}}:Cl_{\text{ex}}:Cl_{\text{eq}}$ as the WURST-CPMG technique is nonquantitative (since each Cl site is likely to have a slightly different T_2 constant) and the relative signal intensity does not accurately reflect the relative populations of the individual chlorine environments. Using results obtained via DFT calculations (vide infra), it is possible to assign the three patterns in the spectrum to the specific chlorine bonding environments. The pattern with the largest absolute magnitude of C_{Q} corresponds to the Cl_{br} environment, consistent with the results obtained for Cp^*ZrCl_3 in the study by Rossini et al.,⁴⁶ but distinct from those for bridging chlorine sites in main group compounds.⁴⁵ This suggests that t[he](#page-16-0) value of C_Q for the bridging chlorine is strongly dependent on the nature of the metal to wh[ich](#page-16-0) it is bonded, as well as the overall structure of the complex. The Cl_{br} has a moderate value of $\eta_{\rm O}$, as is typical for this type of environment. The pattern with the value of η_{Q} close to zero is assigned to the Cl_{ax} , again consistent with high cylindrical symmetry about such terminal M−Cl bonds. The final pattern has a value of $n_{\rm Q}$ closest to unity and is assigned to the Cl_{eq} , which differs from results obtained by Rossini et al. where it was found that terminal chlorine environments had values of $\eta_{\rm Q}$ close to zero. Since the values of $C_{\rm Q}$ for the two types of terminal sites are almost identical, it is clear that η_Q is useful for their differentiation.

In a similar manner, analytical simulations of the entire 35Cl CT pattern, as well as the three contributing patterns, are shown in Figure 5B for TaCl₅. The site assignments are identical to those described for $NbCl₅$. The values for C_O for all three sites of T[a](#page-9-0)Cl₅ are larger than the corresponding values for $NbCl₅$. Since $NbCl₅$ and $TaCl₅$ are isostructural, and tantalum is one row below niobium in the periodic table, it is the relative size of the metal atom which appears to influence the magnitude of the quadrupolar interaction, with the larger central metal atom corresponding to a larger value of C_Q . Interestingly, the M−Cl bond lengths are very similar in these two systems, with the largest difference being 0.03 Å, again discouraging the invoking of simple correlations between bond length and C_Q .

Both of these complexes were previously studied by 35Cl NQR spectroscopy, with the measured quadrupolar frequencies given in Table 2. The room temperature NQR study of $NbCl₅$ reveals seven distinct sites, five of which have comparable NQR frequencies. T[he](#page-4-0) two remaining sites have substantially higher NQR frequencies and are assigned to Cl_{br}, which is consistent with our results.⁷² Similar NQR results were found for $TaCl₅⁸¹$ and are also consistent with this work.

Plane-wave [DF](#page-16-0)T calculations of ³⁵Cl EFG tensors w[ere](#page-16-0) performed on $NbCl₅$ and TaCl₅ (Table 2). Calculated values of C_Q and η_Q distinguish bridging and terminal chlorine sites, not

Figure 6. T_2 relaxation time constants and corresponding CPMG echo trains from experiments on α -WCl₆ at (A) the high-frequency, (B) central, and (C) low-frequency discontinuities.

only by the absolute magnitudes of C_{Q} , but also by their signs: bridging and terminal chlorines have positive and negative values of C_{Q} , respectively. The CASTEP calculations for postgeometry optimized $NbCl₅$ and $TaCl₅$ models are in good agreement with experimental values of C_Q and reasonable agreement for values of $\eta_{\rm Q}$ (Figure 1).

The theoretical ${}^{35}Cl$ EFG tensor orientations were determined for both complexes. Si[n](#page-5-0)ce they are isostructural,

only the results for $NbCl₅$ are shown in Figure 3E. The Cl_{br} sites, which have moderate values of η_{O} , have V_{33} components oriented nearly perpendicular (90.30°) to the M−[C](#page-7-0)l−M plane, which is consistent with observations by Rossini et al.⁴⁶ The moderate values of $\eta_{\rm Q}$ for these sites indicate that the $^{35}\rm Cl$ EFG tensor is non-axial (i.e., $V_{11} \neq V_{22}$), which is consistent [wit](#page-16-0)h the local geometry of a Cl_{br} atom. The Cl_{ax} sites have values of η_{Q} close to zero, with V_{33} components oriented near the directions

^a Core" refers to nonbonding orbitals in the subshells below the valence shell, π LP denotes chlorine lone pairs which may participate in π -bonding, σ LP are lone pairs of σ-symmetry with respect to the Cl–Nb axis, and π , σ, and μ represent orbitals with significant covalent nature. The composition of an LMO with each number signifying percentages. The contributions from each atom as well as the atomic orbital character for selected LMOs are shown. The contributions of each orbital toward V_{11} , V_{22} , and V_{33} are given in au. $\frac{d}{dt}$ The absolute magnitudes of the principal components are reported, as the signs cannot be determined experimentally.

of the M−Cl bonds in each case, as expected. Finally, the Cl_{eq} sites have large values of η_{Q} indicating that V_{11} is the distinct component of the $35C1$ EFG tensor (i.e., the absolute magnitudes of V_{22} and V_{33} are similar). In each system, V_{11} is oriented toward the bridging chlorine environments, and V_{33} is directed near the M−Cl bonds, in good agreement with

³⁵Cl Transverse Relaxation Time (T_2) Measurements. As mentioned above, the ³⁵Cl SSNMR spectra obtained for each of the compounds investigated exhibit variations in spectral intensity that do not precisely match the intensities produced from analytical simulations. For example, higher intensity is consistently observed in the high-frequency (leftmost) regions of the spectra. While there are no detailed studies in the literature that investigate this phenomenon, we have reported instances of this intensity variation in several other papers on ultrawideline NMR of quadrupolar nuclei.^{47,82} We speculate that it may be caused by two factors: (i) tuning limitations of the probe used that may cause uneven excitat[ion/](#page-16-0) detection across the range of frequencies studied and/or (ii) variation of the transverse relaxation times across the breadth of the pattern, which we refer to as T_2 anisotropy.

In order to investigate this further, we measured the transverse relaxation time constants across the patterns by placing the transmitter at frequencies corresponding to different discontinuities of the patterns and acquiring a series of echoes using a CPMG experiment. By plotting the echo intensity against the echo time and fitting the data to an exponential decay, the T_2 values and their associated errors were determined. The process is shown for WCl_6 in Figure 6, and the remaining data are shown in the Supporting Information (Table S8). We found that in most cases, $T₂$ values meas[ur](#page-11-0)ed at the low-frequency discontinuities a[re smaller than those](#page-15-0) measured at higher frequency. The overall differences in T_2 values are not large, certainly well under an order of magnitude. At this time, it is unclear that the tuning characteristics of the probe are responsible for any of the intensity variation, since we

have used the exact same probes and tuning configurations to acquire spectra where the intensity variations do not follow the patterns described above. Clearly, a more detailed study of this intensity variation is required; if T_2 anisotropy is solely responsible for this phenomenon, some interesting physical models and chemical interpretations could arise in the future.

LMO Analysis of the 35 **Cl EFG Tensors.** Upon examining the results obtained for the dimer pentahalide species, certain questions arise: (i) why do the Cl_{br} sites have significantly larger values of C_{O} compared to the terminal chlorines, and (ii) why do the Cl_{eq} sites have values of $\eta_{\rm Q} \approx 1$ when the Cl_{ax} have values of $\eta_{\rm Q} \approx 0$? In order to further probe the origins of the $³⁵$ Cl EFG tensors for the various chlorine environments in the</sup> pentahalide species, we performed localized molecular orbital (LMO) analysis, in which the contributions of each LMO to the ³⁵Cl EFG tensors are determined.⁶⁵ The LMOs are quantum-mechanical representations of bonds and lone pairs as well as core−shell electron pairs. For $Nb₂Cl₁₀$, there are also three-center μ -bonding LMOs centered on the bridging chlorine atoms. Here, we chose to conduct calculations and analysis upon the $Nb₂Cl₁₀$ dimer, which possesses distinct types of terminal and bridging chlorine sites. We note that DFT MO calculations and concomitant LMO analyses were only conducted upon $Nb₂Cl₁₀$ clusters, since these species are sufficiently isolated from the surrounding lattice that 35 Cl EFG tensors are not adversely affected by long-range electrostatic interactions. Compounds like α -WCl₆, β -WCl₆, and MoOCl₄ all feature shorter intermolecular contacts that may influence the 35Cl EFG tensors, so LMO analyses on isolated clusters from these systems were not conducted (a full study of inter- and intramolecular electrostatic effects on EFG tensors for all of these systems is beyond the scope of the current work).

The ³⁵Cl EFG tensor parameters calculated using ADF and CASTEP are in good agreement with one another as well as the experimentally obtained parameters (Table S10, Supporting Information). This indicates that the $35Cl$ EFG tensor parameters are highly dependent upon [EFGs originating within](#page-15-0)

Figure 7. (A) Atom numbering of the Nb₂Cl₁₀ unit used in the LMO analysis. (B) Isosurface representations of the localized LMOs (1) σ (Cl_{ax}− Nb), (2) σ LP Cl_{av} (3) π_x LP Cl_{av} (4) π_y LP Cl_{av} (5) σ(Cl_{eq}–Nb), (6) σ LP Cl_{eq}, (7) π (Cl_{eq}–Nb), (8) π_z LP Cl_{eq}, (9) π_x LP Cl_{eq}, (with some μbonding character), (10) μ (Nb−Cl−Nb), (11) σ LP Cl_{br}, and (12) π _z LP Cl_{br}.

the individual $Nb₂Cl₁₀$ units and almost independent of EFGs arising from intermolecular interactions within the solid. Furthermore, geometry optimization of the structure improves the agreement between the calculated parameters and those obtained experimentally.

The contribution of each LMO to the principal components of the ³⁵Cl EFG tensors are tabulated in Table 3, and the atom labels for the $Nb₂Cl₁₀$ unit are shown in Figure 7A. The individual LMOs that contribute to the EF[Gs](#page-12-0) are shown in Figure 7B as isosurfaces.⁶⁵ The calculations describe contributions from core, lone-pair (LP, with the π and σ referring to local symmetry with res[pe](#page-16-0)ct to the metal-chlorine bond axis), dative Nb−Cl σ-bonds, π-bonds, and three-centered bridging μ bonds. Graphical representations of the ³⁵Cl EFG tensors for the distinct chlorine environments $(Cl_{ax}$, Cl_{aeq} , and Cl_{br}) in the form of polar plots (EFG in the direction of the electric field) are shown in Figure 8. The blue regions of the graphical representations correspond to positive EFGs, and the orange regions represent negative EFGs.

The case of the Cl_{ax} environment is the most straightforward and agrees with previous findings in the literature for terminal chlorines.⁴⁶ The largest component of the EFG tensor, V_{33} is positive and oriented roughly parallel to the Nb−Cl bond axis (Figure [8A](#page-16-0)), indicating a loss of electron density in this direction to the covalent Cl−Nb interactions. The σ (Cl_{ax}−Nb) (MO1) and the σ LP Cl_{ax} (MO2) LMOs both make large negative contributions to the value of V_{33} (Table 3). The π LPs (MO3 and MO4) make large positive contributions that outweigh [th](#page-12-0)ose of the σ orbitals, and therefore the overall V_{33} for the Cl_{ax} is positive. The axial symmetry of the EFG tensor at the Cl_{ax}, with a value of $\eta_{\rm Q}$ close to 0, is due to the cylindrical electronic distribution about the nucleus caused by contributions of MO3 and MO4 to V_{11} and V_{22} , respectively, that are roughly equal in magnitude but opposite in sign.

Like the Cl_{ax}, the Cl_{eq} sites also have a positive V_{33} , which is oriented along the direction of the Nb−Cl bond (Figure 8B).

Figure 8. Graphical representations (polar plots) of 35 Cl EFG tensors of NbCl₅ for (A) Cl_{ax}, (B) Cl_{eq}, and (C) Cl_{br} environments. The blue color indicates a positive EFG while orange indicates a negative EFG. The V_{33} values are +0.71, +0.81, and -1.32 au, for A-C respectively.

The σ (Cl_{eq}–Nb) (MO5) and σ LP Cl_{eq} (MO6) orbitals cause large negative contributions to V_{33} , in a manner similar to the terminal-axial chlorine atoms. The π (Cl_{eq}–Nb) (MO7) and the

Figure 9. Static ³⁵Cl SSNMR spectrum acquired using the WURST-CPMG pulse sequence at 18.8 T for silica-supported WCl₆ with corresponding analytical simulation shown in red.

 π _z LP Cl_{eq} (MO8) orbitals make positive contributions to V_{33} that outweigh those of the σ orbitals and results in an overall positive EFG. The π orbitals of the Cl_{eq} contribute quite differently to the values of V_{11} and V_{22} . MO7 generates a negative contribution to V_{11} and a positive contribution to V_{22} , whereas $MO8$ makes a positive contribution to V_{11} and an even larger negative contribution to V_{22} . The unequal contribution to V_{11} and V_{22} of MO7 and MO8 is reflected in the large asymmetry parameter of the ³⁵Cl EFG tensor ($\eta_{\rm O}$ is near unity). This is contrasted to the case of the Cl_{ax} in which equal involvement from the orbitals with π symmetry leads to an axially symmetric tensor with $\eta_{\rm Q}$ close to zero.

For the Cl_{br} , V_{33} is oriented perpendicular to the equatorial plane formed by the two Nb atoms and the Cl_{br} atom (Figure 8C), as is typical for bridging chlorines.⁴⁷ The Cl_{br} environment consists of two equivalent, weakly covalent Nb-Cl_{br} orbitals [p](#page-13-0)er Cl atom at an angle of 102° and [tw](#page-16-0)o relatively "pure" LP orbitals. The orbitals in the plane of the Nb−Cl−Nb bond (π _x LP Cl_{br} (MO9), $\mu(Nb-Cl-Nb)$ (MO10), and σ LP Cl_{br} $(MO11)$) create positive contributions to V_{33} ; however, these contributions are outweighed by the enormous negative contribution made by the out-of-plane π _z LP Cl_{br} (MO12). MO12 is almost entirely noncovalent in nature and therefore contributes very strongly to the EFG. The result is a large, negative V_{33} at the Cl_{br} nuclei which is oriented in the direction of the lone-pair (MO12). The moderate value of $\eta_{\rm Q}$ for the Cl_{br} nuclei is caused by unequal contributions of the in-plane

orbitals (MO9, MO10, and MO11) to V_{11} and V_{22} .
³⁵Cl SSNMR of WCl₆ Supported on Silica. WCl₆ has been grafted onto a silica support material, as described in the Experimental Section. Our aim in the current work was to see if it was possible to acquire a high quality spectrum of this sample [in a reasonable time f](#page-15-0)rame and make comparisons between this new spectrum and some of the spectra of the tungsten chlorides discussed herein. Only two other attempts at acquiring ³⁵Cl SSNMR spectra of such systems have been reported, by our group (silica-supported $TiCl₄)⁴⁷$ and by Mania et al. (silicasupported $TiCl₄$).²⁶ The ³⁵Cl SSNMR spectrum of this sample, acquired using the WURST-CPMG pulse sequence at a field of 18.8 T, is shown [in](#page-16-0) Figure 9. The spectrum was acquired in a single experiment (i.e., no piecewise acquisitions), over a period of ca. 15.8 h. It is noted that the long experimental times are a result of the low weight percentage of $WCl₆$ loaded onto the silica support material (W wt $% = 5.48\%$). The CT powder pattern spans a range of just over 1.2 MHz, which is far wider than that observed for the spectrum of pure α -WCl₆, which spans only ca. 650 kHz at 21.1 T (it would span ca. 740 kHz at 18.8 T, Figure S20). Simulations of the spectrum of grafted WCl₆ yield C_O = 25.98(5) MHz and η _O = 0.11(1), which are far different [values from](#page-15-0) those of pure WCl_6 ($\text{C}_\text{Q} = 21.0(1) \text{ MHz}$ and $\eta_{Q} = 0.029(5)$). In fact, this new set of quadrupolar parameters bares a striking resemblance to those of $WOCl₄$ (C_O) = 26.05(10), $\eta_{\rm O}$ = 0.115(5)), which suggests that a major change in the structure and bonding of WCl_6 has taken place. A full structural analysis and interpretation of this result is currently underway and beyond the scope of the current work; however, we have again demonstrated that it is possible to acquire high quality ultrawideline $35CI$ SSNMR spectra of diluted, surface-bound species, and that such spectra are potentially very rich in structural information.

■ CONCLUSIONS

Using the WURST-CPMG pulse sequence and the VOCS technique, we have demonstrated that it is possible to acquire broad ³⁵Cl SSNMR powder patterns for diamagnetic chlorinecontaining TM complexes exhibiting a variety of structural motifs. The experiments were conducted at both moderate (9.4 T) and ultrahigh (21.1 T) magnetic field strengths, with the latter affording great reductions in experimental times due to increases in S/N ratios and reductions in CT pattern breadths. The ³⁵Cl SSNMR spectra prove to be very useful as probes of structure, due to the sensitivity of the 35Cl EFG tensors to both subtle and significant differences in molecular geometry and chemical environment.

The ³⁵Cl SSNMR spectra of α -WCl₆ and β -WCl₆ were acquired but were not of high enough resolution to differentiate the two polymorphic forms; however, they could be differentiated by ³⁵Cl NQR experiments. This highlights the efficacy of using SSNMR and NQR in tandem; the rapid acquisition of the SSNMR spectra provides an estimation of the quadrupolar parameters and substantially refines the frequency region in which to search for an NQR signal. The effects of molecular packing in the solid state were also visible in the ³⁵Cl SSNMR spectra of $WOCI_4$ and $MoOCI_4$. The $WOCI_4$ molecules stack in symmetric linear arrays resulting in a single chlorine environment, whereas the $MoOCl₄$ complex forms quasi-dimeric structures resulting in several distinct chlorine environments (i.e., pseudo-bridging and terminal). In the dimeric pentahalide complexes, $NbCl₅$ and $TaCl₅$, the individual chlorine environments were distinguished based on their values of C_Q . It was found that the Cl_{br} environments had larger values of C_{Q} when compared to the terminal chlorine environments. Furthermore, the two terminal chlorine environments, Cl_{ax} and Cl_{eq} , could be distinguished based on their values of η_{Q} , the former having values close to 0, and the latter having values close to unity. It was also observed that the tantalum complex had larger values of C_O for all chlorine environments, suggesting that the size of the metal center influences the value of C_Q .

Plane-wave DFT calculations were completed for each complex. The ³⁵Cl EFG tensor parameters were found to be in generally good agreement with experimental results especially in systems in which intermolecular interactions have little influence on the 35 Cl EFG. The theoretical 35 Cl EFG tensor orientations reveal that V_{33} is directed either along or near the M−Cl bond for chlorine atoms in terminal bonding environments. Conversely, for chlorine atoms in bridging bonding environments, V_{33} 's are oriented approximately perpendicular to the M−Cl−M planes. An LMO analysis completed on the NbCl₅ system revealed that the ³⁵Cl NMR parameters are determined by the composition and relative orientations of the MOs participating in bonding to chlorine. The $\eta_{\rm O}$ value close to zero for the Cl_{ax} arises from equivalent contributions of the π LP orbitals to V_{11} and V_{22} , whereas the π orbitals for the Cl_{eq} do not contribute equivalently, leading to an $\eta_{\rm Q}$ close to unity. The large value of $C_{\rm Q}$ for the Cl_{br} arises from the large, negative contribution of the π _z LP orbital that is perpendicular to the Cl−M−Cl plane, because positive contributions from other orbitals centered on Cl_{br} are reduced due to the true Cl−Nb covalency.

We also demonstrated the application of 35 Cl SSNMR to the study of a transition-metal catalyst that has been grafted onto a silica support material. The resulting spectrum is drastically different than the bulk species, indicating that the structure of the WCl_6 species changes upon grafting. Given the rapidity with which the ³⁵Cl SSNMR spectra can be acquired, and the fact that "focused" NQR experiments can be conducted in the wake of SSNMR experiments to examine site specificities, we believe that an enormous number of chlorinated transitionmetal complexes can be structurally probed in both the bulk and supported forms. We hope to establish this protocol as a strong basis for the study of chlorine-containing transitionmetal complexes and extend this study to catalytic systems of increasing complexity, with the aim of studying immobilized heterogeneous catalysts with very dilute chlorine contents.

■ ASSOCIATED CONTENT

9 Supporting Information

Details on NMR acquisition parameters, pXRD patterns, additional ³⁵Cl SSNMR data, details on computational chemistry and LMO analysis, and analytical simulations of ³⁵Cl SSNMR spectra. This material is available free of charge via the Internet at http://pubs.acs.org.

■ AUTHOR IN[FORMATION](http://pubs.acs.org)

Corresponding Author

*Phone: (519) 253-3000 x3548. Fax: (519) 973-7098. E-mail: rschurko@uwindsor.ca.

Notes

[The authors declare no](mailto:rschurko@uwindsor.ca) competing financial interest.

■ ACKNOWLEDGMENTS

R.W.S. thanks NSERC for funding this research in the form of a Discovery Grant and Discovery Accelerator Supplement. R.W.S. is also grateful for an Early Researcher Award from the Ontario Ministry of Research and Innovation. We are also grateful for the funding of the Laboratories for Solid-State Characterization at the University of Windsor from the Canadian Foundation for Innovation, the Ontario Innovation Trust, and the University of Windsor. This work was made possible, in part, by the facilities of the Shared Hierarchical Academic Research Computing Network (SHARCNET: http://www.sharcnet.ca). We are grateful to Dr. Victor Terskikh at the National Ultrahigh-field NMR Facility for [Solids in Ottawa, Ontario](http://www.sharcnet.ca), Canada (http://nmr900.ca/), as well as Dr. Kristopher Harris, for acquiring ³⁵Cl SSNMR data at 21.1 T. J.A. acknowledges support fr[om the National](http://nmr900.ca/) Science Foundation (CHE-1265833).

■ REFERENCES

(1) Ajiboye, S. I.; Iqbal, J.; Peacock, R. D.; Prouff, N.; Taylor, G. J.; Winfield, J. M.; Liu, X. Fluor. Chem. 1998, 91, 213−218.

(2) Englert, U.; Pampaloni, G.; Puccini, F.; Volpe, M. J. Organomet. Chem. 2007, 692, 3144−3150.

(3) Sim, A.; Cant, N. W.; Trimm, D. L. Int. J. Hydrogen Energy 2010, 35, 8953−8961.

(4) Nguyen, A. I.; Blackmore, K. J.; Carter, S. M.; Zarkesh, R. A.; Heyduk, A. F. J. Am. Chem. Soc. 2009, 131, 3307−3316.

(5) Mokdsi, G.; Harding, M. M. J. Inorg. Biochem. 2001, 86, 611−616. (6) Waern, J. B.; Harris, H. H.; Lai, B.; Cai, Z.; Harding, M. M.;

Dillon, C. T. J. Biol. Inorg. Chem. 2005, 10, 443−452.

(7) Kö pf-Maier, P.; Klapötke, T. J. Cancer Res. Clin. Oncol. 1992, 118, 216−221.

(8) Matthias, B. T.; Geballe, T. H.; Geller, S.; Corenzwit, E. Phys. Rev. 1954, 95, 1435.

(9) Raub, C. J.; Sweedler, R.; Jensen, M. A.; Broadston, S.; Matthiasg, T. Phys. Rev. Lett. 1964, 13, 746−747.

(10) Bukowski, Z.; Badurski, D.; Stepien-Damm, J.; Troc, R. Solid State Commun. 2002, 123, 283−286.

(11) Nowak, I.; Ziolek, M. Chem. Rev. 1999, 99, 3603−3624.

(12) Tanabe, K.; Okazaki, S. Appl. Catal., A 1995, 133, 191−218.

(13) Wallace, K. C.; Liu, A. H.; Davis, W. M.; Schrock, R. R. Organomet. 1989, 8, 644−654.

(14) Ushikubo, T.; Wada, K. J. Catal. 1994, 148, 138−148.

(15) Schrock, R. R.; Hoveyda, A. H. Angew. Chem., Int. Ed. 2003, 42, 4592−4633.

(16) Yandulov, D. V.; Schrock, R. R. Science 2003, 301, 76−78.

(17) Balcar, H.; Čejka, J.; Sedláček, J.; Svoboda, J.; Zedník, J.; Bastl,

Z.; Bosáček, V.; Vohlídal, J. J. Mol. Catal. A 2003, 203, 287–298.

(18) Garrett, C. E.; Prasad, K. Adv. Synth. Catal. 2004, 346, 889−900.

Inorganic Chemistry Article

- (19) Collman, J. P.; Hegedu, L. S.; Cooke, M. P.; Norton, J. R.; Dolcetti, G.; Marquardt, D. N. J. Am. Chem. Soc. 1972, 94, 1789−1790.
- (20) Dumont, W.; Poulin, J.; Dang, T.; Kagan, H. B. J. Am. Chem. Soc. 1973, 95, 8295−8299.
- (21) Maschmeyer, T.; Rey, F.; Sankar, G.; Thomas, J. M. Nature 1995, 378, 159−162.
- (22) Eisen, M. S.; Marks, T. J. J. Am. Chem. Soc. 1992, 114, 10358− 10368.
- (23) Joubert, J.; Delbecq, F.; Sautet, P.; Le Roux, E.; Taoufik, M.; Thieuleux, C.; Blanc, F.; Copéret, C.; Thivolle-Cazat, J.; Basset, J.-M. J. Am. Chem. Soc. 2006, 128, 9157-9169.
- (24) Blanc, F.; Thivolle-Cazat, J.; Basset, J.-M.; Copéret, C.; Hock, A. S.; Tonzetich, Z. J.; Sinha, A.; Schrock, R. R. J. Am. Chem. Soc. 2007, 129, 1044−1045.
- (25) Blanc, F.; Basset, J.-M.; Coperet, C.; Sinha, A.; Tonzetich, Z. J.; ́ Schrock, R. R.; Solans-Monfort, X.; Clot, E.; Eisenstein, O.; Lesage, A.; Emsley, L. J. Am. Chem. Soc. 2008, 130, 5886−5900.
- (26) Mania, P.; Verel, R.; Jenny, F.; Hammond, C.; Hermans, I. Chem.Eur. J. 2013, 19, 9849−9858.
- (27) Blanc, F.; Copéret, C.; Lesage, A.; Emsley, L. Chem. Soc. Rev. 2008, 37, 518−526.
- (28) Zhang, W.; Xu, S.; Han, X.; Bao, X. Chem. Soc. Rev. 2012, 41, 192−210.
- (29) Jezequel, M.; Dufaud, V.; Ruiz-Garcia, M. J.; Carrillo-Hermosilla, F.; Neugebauer, U.; Niccolai, G. P.; Lefebvre, F.; Bayard, F.; Corker, J.; Fiddy, S.; Evans, J.; Broyer, J. P.; Malinge, J.; Basset, J. M. J. Am. Chem. Soc. 2001, 123, 3520−3540.
- (30) Ahn, H.; Marks, T. J. J. Am. Chem. Soc. 2002, 124, 7103−7110. (31) Le Roux, E.; Chabanas, M.; Baudouin, A.; de Mallmann, A.; Coperet, C.; Quadrelli, E. A.; Thivolle-Cazat, J.; Basset, J.; Lukens, W.;
- Lesage, A.; Emsley, L. J. Am. Chem. Soc. 2004, 126, 13391−13399.
- (32) Hung, I.; Schurko, R. W. J. Phys. Chem. B 2004, 108, 9060− 9069.
- (33) Lo, A. Y. H.; Bitterwolf, T. E.; Macdonald, C. L. B.; Schurko, R. W. J. Phys. Chem. A 2005, 109, 7073-7087.
- (34) Hamaed, H.; Lo, A. Y. H.; Lee, D. S.; Evans, W. J.; Schurko, R. W. J. Am. Chem. Soc. 2006, 128, 12638-12639.
- (35) Rossini, A. J.; Hung, I.; Schurko, R. W. J. Phys. Chem. Lett. 2010, 1, 2989−2998.
- (36) Rossini, A. J.; Hung, I.; Johnson, S. A.; Slebodnick, C.; Mensch, M.; Deck, P. A.; Schurko, R. W. J. Am. Chem. Soc. 2010, 132, 18301− 18317.
- (37) Larsen, F. H.; Jakobsen, H. J.; Ellis, P. D.; Nielsen, N. C. J. Magn. Reson. 1998, 131, 144−147.
- (38) O'Dell, L. A.; Schurko, R. W. Chem. Phys. Lett. 2008, 464, 117− 130.
- (39) Harris, K. J.; Lupulescu, A.; Lucier, B. E. G.; Frydman, L.; Schurko, R. W. J. Magn. Reson. 2012, 224, 38−47.
- (40) Harris, R. K.; Becker, E. D.; Menezes, S. M. C. D. E. Pure Appl. Chem. 2001, 73, 1795−1818.
- (41) Kentgens, A. P. M. Geoderma 1997, 80, 271−306.
- (42) Chapman, R. P.; Hiscock, J. R.; Gale, P. A.; Bryce, D. L. Can. J. Chem. 2011, 89, 822−834.
- (43) Bryce, D. L.; Gee, M.; Wasylishen, R. E. J. Phys. Chem. A 2001, 105, 10413−10421.
- (44) Hamaed, H.; Pawlowski, J. M.; Cooper, B. F. T.; Fu, R.; Eichhorn, S. H.; Schurko, R. W. J. Am. Chem. Soc. 2008, 130, 11056− 11065.
- (45) Chapman, R. P.; Bryce, D. L. Phys. Chem. Chem. Phys. 2009, 11, 6987−6998.
- (46) Rossini, A. J.; Mills, R. W.; Briscoe, G. A.; Norton, E. L.; Geier, S. J.; Hung, I.; Zheng, S.; Autschbach, J.; Schurko, R. W. J. Am. Chem. Soc. 2009, 131, 3317−3330.
- (47) Johnston, K. E.; O'Keefe, C. A.; Gauvin, R. M.; Trebosc, J.; ́ Delevoye, L.; Amoureux, J.-P.; Popoff, N.; Taoufik, M.; Oudatchin, K.; Schurko, R. W. Chem.-Eur. J. 2013, 19, 12396-12414.
- (48) Perras, F. A.; Bryce, D. L. Angew. Chem., Int. Ed. Engl. 2012, 51, 4227−4230.
- (49) Schurko, R. W. In Encyclopedia of Magnetic Resonance; John Wiley and Sons: New York, 2012; pp 77−94.
- (50) Schurko, R. W. Acc. Chem. Res. 2013, 46, 1985−1995.
- (51) O'Dell, L. A.; Rossini, A. J.; Schurko, R. W. Chem. Phys. Lett. 2009, 468, 330−335.
- (52) Taylor, J. C.; Wilson, P. W. Acta Crystallogr. Sect. B Struct. Sci. 1974, 30, 1216−1220.
- (53) Popoff, N.; Espinas, J.; Goure, E.; Boyron, O.; Le Roux, E.; ́ Basset, J.-M.; Gauvin, R. M.; De Mallmann, A.; Taoufik, M. Macromol. Rapid Commun. 2011, 32, 1921−1924.
- (54) KrausW.; Nolze, G.PowderCell for Windows, 2000.
- (55) Massiot, D.; Farnan, I.; Gautier, N.; Trumeau, D.; Trokiner, A.; Coutures, J. P. Solid State Nucl. Magn. Reson. 1995, 4, 241−248.
- (56) Medek, A.; Frydman, V.; Frydman, L. J. Phys. Chem. A 1999, 103, 4830−4835.
- (57) Tang, J. A.; Masuda, J. D.; Boyle, T. J.; Schurko, R. W. ChemPhysChem 2006, 7, 117−130.
- (58) Eichele, K.; Wasylishen, R. E.WSolids: Solid-State NMR Spectrum Simulation Package, 2001.
- (59) Perras, F. A.; Widdifield, C. M.; Bryce, D. L. Solid State Nucl. Magn. Reson. 2012, 45−46, 36−44.
- (60) Clark, S. J.; Segall, M. D.; Pickard, C. J.; Hasnip, P. J.; Probert,
- M. I. J.; Refson, K.; Payne, M. C. Z. Kristallogr. 2005, 220, 567−570.
- (61) Baerends, E. J. Z. T.; Autschbach, J. Amsterdam Density Functional.
- (62) Van Lenthe, E.; Snijders, J. G.; Baerends, E. J. J. Chem. Phys. 1996, 105, 6505−6516.
- (63) Hammer, B.; Hansen, L.; Nørskov, J. Phys. Rev. B 1999, 59, 7413−7421.
- (64) Perdew, J.; Burke, K.; Ernzerhof, M. Phys. Rev. Lett. 1996, 77, 3865−3868.
- (65) Autschbach, J.; Zheng, S.; Schurko, R. W. Concepts Magn. Reson. 2010, 36A, 84−126.
- (66) Reed, A. E.; Weinhold, F. J. Chem. Phys. 1985, 83, 1736.
- (67) Glendening, E. D.; Badenhoop, J. K.; Reed, A. E.; Carpenter, J.
- E.; Bohmann, J. A.; Morales, C. M.; Weinhold, F. NBO 5.0, 2001.
- (68) Harris, R. K. Analyst 2006, 131, 351−373.
- (69) Lucken, E. A. C. In Nuclear Quadrupole Coupling Constants; Academic Press: New York, 1969; pp 34−45.
- (70) Das, T. P.; Hahn, E. L. In Nuclear Quadrupole Resonance Spectroscopy; Academic Press: New York, 1958; pp 3−7.
- (71) Smith, D. K.; Landingham, R. L.; Smith, G. S.; Johnson, Q. Acta Crystallogr. Sect. B Struct. Sci. 1968, 24, 1563.
- (72) Pisarev, E. A.; Semin, G. K.; Drobot, D. V.; Kuznetsov, S. I.; Bryukhova, E. V. Zh. Neorg. Khim. 1974, 23, 1171−1182.
- (73) Suits, B. H. Nuclear Quadrupolar Spectroscopy in Handbook of Applied Solid-State Spectroscopy; Vij, D. R., Ed.; Springer: New York, 2006.
- (74) Widdifield, C. M.; Chapman, R. P.; Bryce, D. L. Annu. Rep. NMR Spectrosc. 2009, 66, 195.
- (75) Rossini, A. J.; Hamaed, H.; Schurko, R. W. J. Magn. Reson. 2010, 206, 32−40.
- (76) Hess, H.; Hartung, H. Z. Anorg. Chem. 1966, 344, 157.
- (77) Taylor, J. C.; Waugh, A. B. Dalton Trans. 1980, 10, 2006−2009.
- (78) Lucken, E. A. C. In Nuclear Quadrupole Coupling Constants;
- Academic Press: New York, 1969; pp 120−130.
- (79) Zalkin, A.; Sands, D. E. Acta Crystallogr. 1958, 11, 615−619.
- (80) Rabe, S.; Mueller, U. Z. Krist. New Cryst. Struct. 2000, 215, 1− 2.
- (81) Semin, G. K.; Kuznetsov, S. I.; Alimov, I. M.; Khotsianova, T. L.; Bryukhova, E. V.; Nisselson, L. A.; Tretyakova, K. V. Inorg. Chim. Acta 1974, 13, 181.
- (82) Lucier, B. E. G.; Tang, J. A.; Schurko, R. W.; Bowmaker, G. A.; Healy, P. C.; Hanna, J. V. J. Phys. Chem. C 2010, 114, 7949−7962.
- (83) Dye, J. L.; Ellaboudy, A. S.; Kim, J. Solid State NMR of Quadrupolar Nuclei; Marcel Dekker, Inc.: New York, 1991.