ChemComm



COMMUNICATION

View Article Online



Cite this: Chem. Commun., 2015 51 4993

Received 27th January 2015, Accepted 17th February 2015

DOI: 10.1039/c5cc00499c

www.rsc.org/chemcomm

Synthesis and characterisation of an open-cage fullerene encapsulating hydrogen fluoride†

Andrea Krachmalnicoff, Richard Bounds, Salvatore Mamone, Malcolm H. Levitt. Marina Carravetta and Richard J. Whitby*

The first encapsulation of hydrogen fluoride in an open-cage fullerene is reported. Solution and solid-state NMR spectra of the novel open-cage endofullerene are described.

Open-cage fullerenes provide the opportunity of trapping atoms and molecules inside an inert three-dimensional environment - a molecular-scale "nanolaboratory". 1-3 Their supramolecular structure makes the study of effectively isolated molecules practically feasible over a wide range of conditions.4 Physical phenomena such as spin-isomer conversion have been observed inside fullerene cages.⁵ The incorporation of the small molecules H₂,⁶⁻¹⁰ $H_2O_{,10-15}^{10-15}N_{2,1}^{15,16}CO_{,16,17}^{16,17}NH_{3,18}^{18}$ and $CH_{4,19}^{19}$ inside the cavity of open-cage fullerenes has been reported. Completion of "Molecular Surgery" to reform the pristine C60 cage has been achieved only for $H_2@C_{60}$, 7,10 and $H_2O@C_{60}$. 10,14 Hydrogen fluoride (HF) is one of the most studied molecules, both theoretically and experimentally.²⁰ Empirical studies on isolated HF molecules are complicated by its high chemical reactivity and its marked tendency to form aggregates and strong hydrogen-bonds to Lewis bases. Many experiments show the constant presence of oligomers, even in the gas phase.21 Enclosure in a fullerene would provide a constrained non-coordinating environment for a single molecule of HF and potentially allow novel studies on its spectral properties, particularly at cryogenic temperatures. Although the fully enclosed and symmetric HF@C₆₀ structure would be ideal, open-cage fullerene hosts could provide many of the same advantages. Important physical properties such as ferroelectricity have been anticipated for endofullerenes enclosing freely-rotating molecules with an electric dipole moment.²²

Herein we describe the insertion of HF into the cavity of open-cage fullerene 1, and the solution and solid-state NMR study of endofullerene HF(a)1.

Chemistry, University of Southampton, Southampton SO17 1BJ, UK. E-mail: rjw1@soton.ac.uk

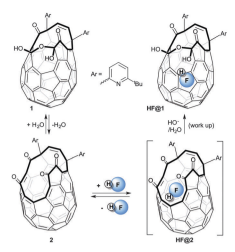
Density Functional Theory (DFT) calculations gave activation energies of 64.3, 52.2. and 29.8 kJ mol⁻¹ respectively for the entry of H₂, H₂O and HF into 2.²³ The lower barrier for the incorporation of H₂O compared to H₂, though surprising, agrees with the experimental evidence¹⁰ (temperatures of 100 and 120 °C are required respectively) and is presumably due to attractive dipolar interactions between the polar molecule and the open fullerene neck. The very low activation energy predicted for the entry of HF into 2 and the possibility of trapping the endohedral HF via hemiacetal formation made HF@1 a viable target. The calculated binding energy for HF inside 2 (25.9 kJ mol⁻¹) and the barrier for the release of HF from HF(a)2 (55.7 kJ mol⁻¹) suggested that the loss of HF from HF@2 would be slow at room temperature allowing HF@1 to form.

The above calculations prompted us to attempt the filling of 2 with HF. Gaseous hydrogen fluoride forms relatively stable polymeric adducts with several organic bases;²⁴ therefore we selected such compounds as a convenient source of anhydrous hydrogen fluoride. When a solution of tetra-ketone 2 or of its hydrate 1 was treated with a large excess of 70% w/w hydrogen fluoride in pyridine (HF-Py), HF@1 was isolated after basic work-up and chromatography (Scheme 1).

The filling factor of HF@1 was established by ¹H NMR spectroscopy, comparing the integral values of the endohedral HF proton with the protons on the exohedral groups. The highest filling factor (50%) was achieved by equilibrating a solution of 1 or 2 in dichloromethane with an excess (200 eq. HF) of HF-Py at room temperature. In these conditions the equilibrium was reached within 24 hours and prolonged reaction time did not afford a higher filling factor; 50% filled HF@1 was isolated after work up and chromatography in 89% yield. A larger excess (300 eq. HF) of HF-pyr did not increase the filling factor, but a lower excess (100 eq. HF) gave only 40% filled HF@1. Both compounds 1 and 2 can be used as substrate, as the acidic reaction medium is evidently able to afford the dehydration of 1 to form 2. The filling at 4 °C proceeded at a much slower rate (~6 days) and did not improve the filling factor. At 80 °C a lower filling factor was obtained, as would be expected for an entropically disfavoured

[†] Electronic supplementary information (ESI) available: Experimental procedures. compound characterization data and details of calculations. See DOI: 10.1039/

Communication ChemComm



Scheme 1 Generation of HF@1 from compound 1

process, and decomposition of the substrate occurred giving 30% filled HF@1 in slightly lower isolated yield (80%). When the filling was carried out in non-chlorinated solvents such as benzene a biphasic system was formed and the substrate was quantitatively extracted into the HF-Py layer. After work-up and chromatography, 50% filled HF@1 was isolated in very good yield (88%). The ESI + MS spectrum of the isolated compounds displays signals at m/z 1121 and 1141 respectively for the molecular ions $[1 + H]^+$ and $[HF(2)1 + H]^+$.

Solution state NMR typically reveals unusual chemical shifts for the endohedral nuclei due to the strong magnetic shielding effect of the fullerene cage. 4,25 Indeed the ¹H signal from the HF molecule in **HF(a)1** appears as a doublet centred at $\delta = -6.55$ ppm with a $I_{\rm HF}$ of 508 Hz (Fig. 1). This chemical shift is similar to those reported for the endohedral protons of $H_2@1$ and $H_2O@1$ which resonate at δ -7.17 and -9.84 ppm respectively. The large $J_{\rm HF}$ value is comparable to that reported for HF in the gas phase.²⁶

A doublet with a $J_{\rm HF}$ of 508 Hz is present in the ¹⁹F NMR at δ -223.91 ppm (Fig. 2); the two lines coalesce into a singlet in the proton decoupled spectrum.

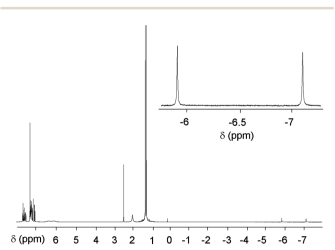


Fig. 1 1 H NMR spectrum (400 MHz, CDCl₃) of **HF@1**. The δ scale is referenced to CHCl₃.

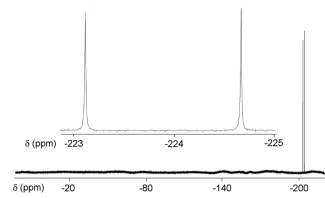


Fig. 2 $\,^{19}\text{F}$ NMR (376 MHz, CDCl}3) spectrum of compound HF@1. The δ scale is referenced to CFCl3

Solid-state NMR has been used for the characterization of molecular endofullerenes.^{25,27,28} The benefits of solid-state NMR include a much larger temperature range, even descending into the cryogenic regime⁵ and also the preservation of anisotropic interactions, which are averaged out in solution state NMR, but which contain information on the local structure, dynamics and symmetry. In the case of HF(a)1, the relevant anisotropic nuclear interactions include the chemical shift anisotropy (CSA), which reflects the electronic environment, and the dipole-dipole interaction between the ¹H and ¹⁹F nuclei of the endohedral molecule.

Magic angle spinning (MAS) may be used to partly average out anisotropic interactions and produce high resolution spectra, ²⁹ which may still reveal the presence of significant anisotropies through spinning sidebands, which appear at multiples of the spinning frequency, and whose intensities are characteristic of the anisotropic interaction parameters.

The ¹H solid-state MAS spectrum (Fig. 3), collected in a magnetic field of 19.96 T (850 MHz for ¹H) at a spinning frequency of 20 kHz, displays an intense signal arising from the numerous exohedral protons and narrow peaks from the endohedral HF, reflecting the relatively isolated magnetic environment of the endohedral

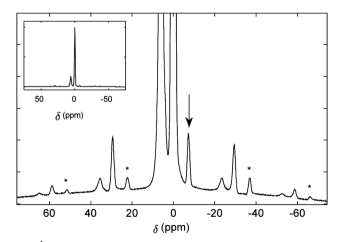


Fig. 3 ¹H NMR spectrum of **HF@1** recorded at 850 MHz (19.96 T) and a temperature of 263 K. The spinning frequency was 20 kHz. The -7 ppm centre band of the HF resonance is shown by the arrow. The spinning side bands are marked with asterisks.

ChemComm Communication

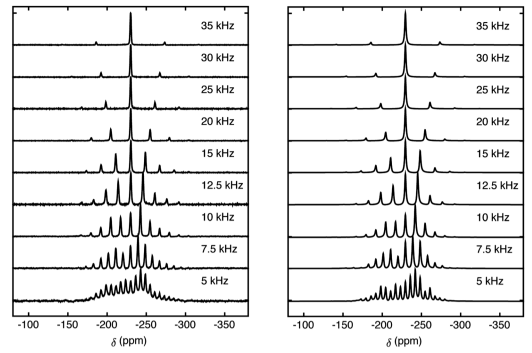


Fig. 4 (left) 19 F spectra of **HF@1** recorded at 19.96 T and a temperature of 263 K, at the indicated spinning frequencies. The 19 F chemical shift scale is referenced to β-polyvinylidene fluoride (PVDF); (right) simulated spectra with parameters determined from best fit to experiment. We used SPINACH³⁰ and SIMPSON³¹ for simulations. Details of simulations can be found in ESI.†

molecule and its rapid molecular motion inside the cage. This is in agreement with studies on other open-cage endofullerenes.²⁸

The 1 H chemical shift is -7 ppm in the solid-state NMR spectrum, which is similar to the liquid-state value. The $J_{\rm HF}$ coupling is not resolved in the solid-state NMR spectrum. Spinning sidebands can be seen at multiples of the spinning frequency, 20 kHz.

Magic-angle-spinning 19 F spectra of **HF@1** are shown in Fig. 4 for a set of spinning speeds. The 19 F isotropic chemical shift is observed at -230 ppm.

There are two anisotropic interactions that contribute to the generation of sidebands at low spinning frequencies, namely the dipole–dipole $^1\text{H}^{-19}\text{F}$ interaction and the ^{19}F CSA. Numerical simulations of the NMR spectra were generated for combinations of isotropic chemical shift, asymmetry parameter, chemical shift anisotropy and the dipolar interaction using both SPINACH³⁰ and SIMPSON.³¹ The direct dipolar interaction was determined to be -7.5 ± 2.5 kHz and the ^{19}F chemical shift anisotropy as 44.1 ppm \pm 1.7 ppm, with the biaxiality determined to be 0.6 \pm 0.05. The ^{19}F CSA tensor is found to be orientated at (α,β,γ) = $(0,90^{\circ},0)$ with respect to the dipolar coupling tensor.

The direct dipolar coupling constant for an immobile $^{1}\text{H}-^{19}\text{F}$ pair separated by 0.91 Å would be -150 kHz. The small value of the observed dipolar coupling therefore indicates that the HF molecule rotates rapidly and almost isotropically inside the open fullerene cage. The second-rank order parameter of the HF molecule, indicating the degree of anisotropy of its rotational motion, is only $\sim 5\%$.

In this article we report the first HF-endofullerene and its study by solution and solid-state NMR. The solution NMR data

shows a large $^1\text{H}^{-19}\text{F}$ *J*-coupling of 508 Hz, similar to that obtained for HF in the gas phase. The solid-state NMR spectra indicate that the HF molecule rotates rapidly and almost isotropically in the supramolecular complex. Unfortunately attempts to close the cage to give HF@C60 as reported for H2O@C60¹⁰ resulted in complete loss of HF.

This research was supported by the EPSRC (EP/I029451/1), the European Regional Development Fund (ERDF) Interreg-IVB, MEET project and the ERC. MC acknowledges the Royal Society for her University Research Fellowship. We are grateful for EPSRC Core Capability Funding (EP/K039466/1). We acknowledge the use of the IRIDIS High Performance Computing Facility and associated support services at the University of Southampton. The UK 850 MHz solid-state NMR Facility used in this research was funded by EPSRC and BBSRC, as well as the University of Warwick including *via* part funding through Birmingham Science City Advanced Materials Projects 1 and 2 supported by Advantage West Midlands (AWM) and the ERDF. We would like to thank Dinu Iuga for experimental support at the 850 MHz NMR facility and Ilya Kuprov for help with SPINACH.

Notes and references

- 1 M. H. Levitt and A. J. Horsewill, *Philos. Trans. R. Soc. London, Ser. A*, 2013, 371, 20130124.
- (a) L. Gan, D. Yang, Q. Zhang and H. Huang, *Adv. Mater.*, 2010, 22, 1498;
 (b) G. C. Vougioukalakis, M. M. Roubelakis and M. Orfanopoulos, *Chem. Soc. Rev.*, 2010, 39, 817.
- 3 S. Iwamatsu and S. Murata, Synlett, 2005, 2117.
- 4 M. H. Levitt, Philos. Trans. R. Soc. London, Ser. A, 2013, 371, 20120429.
- 5 (a) N. J. Turro, A. A. Marti, J. Y. C. Chen, S. Jockusch, R. G. Lawler, M. Ruzzi, E. Sartori, S. C. Chuang, K. Komatsu and Y. Murata, *J. Am. Chem. Soc.*, 2008, **130**, 10506; (b) Y. Li, X. Lei, S. Jockusch, J. Y. C. Chen,

M. Frunzi, J. A. Johnson, R. G. Lawler, Y. Murata, M. Murata, K. Komatsu and N. J. Turro, *J. Am. Chem. Soc.*, 2010, 132, 4042; (c) C. Beduz, M. Carravetta, J. Y.-C. Chen, M. Concistré, M. Denning, M. Frunzi, A. J. Horsewill, O. G. Johannessen, R. Lawler, X. Lei, M. H. Levitt, Y. Li, S. Mamone, Y. Murata, U. Nagel, T. Nishida, J. Ollivier, S. Rols, T. Rõõm, R. Sarkar, N. J. Turro and Y. Yang, *Proc. Natl. Acad. Sci. U. S. A.*, 2012, 109, 12894; (d) S. Mamone, M. Concistre, E. Carignani, B. Meier, A. Krachmalnicoff, O. G. Johannessen, X. Lei, Y. Li, M. Denning, M. Carravetta, K. Goh, A. J. Horsewill, R. J. Whitby and M. H. Levitt,

Communication

J. Chem. Phys., 2014, 140, 194306.

- 6 Y. Rubin, T. Jarrosson, G.-W. Wang, M. D. Bartberger, K. N. Houk, G. Schick, M. Saunders and R. J. Cross, Angew. Chem., Int. Ed., 2001, 40, 1543.
- 7 K. Komatsu, M. Murata and Y. Murata, Science, 2005, 307, 238.
- 8 S. Iwamatsu, S. Murata, Y. Andoh, M. Minoura, K. Kobayashi, N. Mizorogi and S. Nagase, *J. Org. Chem.*, 2005, **70**, 4820.
- 9 Y. Hashikawa, M. Murata, A. Wakamiya and Y. Murata, Org. Lett., 2014, 16, 2970.
- 10 A. Krachmalnicoff, M. H. Levitt and R. J. Whitby, *Chem. Commun.*, 2014, 50, 13037.
- 11 S. Iwamatsu, T. Uozaki, K. Kobayashi, S. Re, S. Nagase and S. Murata, J. Am. Chem. Soc., 2004, 126, 2668.
- 12 Z. Xiao, J. Yao, D. Yang, F. Wang, S. Huang, L. Gan, Z. Jia, Z. Jiang, X. Yang, B. Zheng, G. Yuan, S. Zhang and Z. Wang, *J. Am. Chem. Soc.*, 2007, 129, 16149.
- 13 Q. Zhang, T. Pankewitz, S. Liu, W. Klopper and L. Gan, Angew. Chem., Int. Ed., 2010, 49, 9935.
- 14 K. Kurotobi and Y. Murata, Science, 2011, 333, 613.
- 15 T. Futagoishi, M. Murata, A. Wakamiya, T. Sasamori and Y. Murata, Org. Lett., 2013, 15, 2750.
- 16 C. M. Stanisky, R. J. Cross and M. Saunders, J. Am. Chem. Soc., 2009, 131, 3392.
- 17 S. Iwamatsu, C. M. Stanisky, R. J. Cross, M. Saunders, N. Mizorogi, S. Nagase and S. Murata, *Angew. Chem., Int. Ed.*, 2006, 45, 5337.
- 18 K. E. Whitener Jr., M. Frunzi, S. Iwamatsu, S. Murata, R. J. Cross and M. Saunders, *J. Am. Chem. Soc.*, 2008, **130**, 13996.

- 19 K. E. Whitener Jr., R. J. Cross, M. Saunders, S. Iwamatsu, S. Murata, N. Mizorogi and S. Nagase, J. Am. Chem. Soc., 2009, 131, 6338.
- 20 D. Feller and K. A. Peterson, THEOCHEM, 1997, 400, 69.
- (a) R. L. Jarry and W. Davis, J. Phys. Chem., 1953, 57, 600;
 (b) D. F. Smith, J. Chem. Phys., 1958, 28, 1040;
 (c) R. A. Oriani and C. P. Smyth, J. Am. Chem. Soc., 1948, 70, 125.
- 22 J. Cioslowski and A. Nanayakkara, Phys. Rev. Lett., 1992, 69, 2871.
- 23 DFT calculations were carried out using MO62-X/cc-pVDZ at B3LYP/6-31G(d) geometries. Details in the ESI†.
- 24 (a) G. A. Olah, J. T. Welch, Y. D. Vankar, M. Nojima, I. Kerekes and J. A. Olah, J. Org. Chem., 1979, 44, 3872; (b) G. A. Olah, T. Mathew, A. Goeppert, B. Torok, I. Bucsi, X. Y. Li, Q. Wang, E. R. Marinez, P. Batamack, R. Aniszfeld and G. K. Prakash, J. Am. Chem. Soc., 2005, 127, 5964.
- 25 M. Carravetta, A. Danquigny, S. Mamone, F. Cuda, O. G. Johannessen, I. Heinmaa, K. Panesar, R. Stern, M. C. Grossel, A. J. Horsewill, A. Samoson, M. Murata, Y. Murata, K. Komatsu and M. H. Levitt, *Phys. Chem. Chem. Phys.*, 2007, 9, 4879.
- (a) C. MacLean and E. L. Mackor, J. Chem. Phys., 1961, 34, 2207;
 (b) J. S. Muenter, J. Chem. Phys., 1970, 52, 6033; (c) J. S. Martin and F. Y. Fujiwara, J. Am. Chem. Soc., 1974, 96, 7632.
- 27 M. Concistrè, S. Mamone, M. Denning, G. Pileio, X. Lei, Y. Li, M. Carravetta, N. J. Turro and M. H. Levitt, *Philos. Trans. R. Soc. London, Ser. A*, 2013, 371, 20120102.
- 28 M. Carravetta, Y. Murata, M. Murata, I. Heinmaa, R. Stern, A. Tontcheva, A. Samoson, Y. Rubin, K. Komatsu and M. H. Levitt, J. Am. Chem. Soc., 2004, 126, 4092.
- 29 M. H. Levitt, Spin Dynamics: Basics of Nuclear Magnetic Resonance, Wiley, 2nd edn, 2008.
- 30 (a) H. J. Hogben, M. Krzystyniak, G. T. P. Charnock, P. J. Hore and I. Kuprov, J. Magn. Reson., 2011, 208, 179; (b) L. J. Edwards, D. V. Savostyanov, A. A. Nevzorov, M. Concistrè, G. Pileio and I. Kuprov, J. Magn. Reson., 2013, 235, 121.
- 31 M. Bak, J. T. Rasmussen and N. C. Nielsen, J. Magn. Reson., 2000, 147, 296.