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Published in:
Physical Review B

Citation for published version (APA):

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Electronic structure of pristine and intercalated \ce{Sc_3N@C_{80}} metallofullerene


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We report a study of the electronic structure and charge transfer in the metallofullerene \ce{Sc_3N@C_{80}} using photoemission and x-ray absorption spectroscopy. Through a comparison of the x-ray absorption spectrum of \ce{Sc_3N@C_{80}} at the Sc L\textsubscript{3,2} edge with atomic multiplet calculations, the Sc 3\textit{d} electron count is determined to be 0.6, thus giving an effective Sc valency of 2.4. With the N atom gaining a full electronic shell by means of covalent bonding with the Sc (also involving the Sc 3\textit{d} electron density observed in the x-ray absorption experiments), the remaining six valence electrons of the \ce{Sc_3N} cluster are then transferred to the carbon cage which stabilizes the \ce{C_{80}} cage structure with \textit{I}\textsubscript{h} symmetry, a structure which is not energetically favored in neutral \ce{C_{80}}. The presence of the highly symmetric \textit{I}\textsubscript{h} cage structure is further supported by the observation of distinct fine structure in the valence band photoemission spectra of the endohedral, which results from the high degree of effective degeneracy of the electronic states in the molecule. Finally, the results of investigations of K-doped \ce{Sc_3N@C_{80}} using photoemission give insight into the K,\ce{Sc_3N@C_{80}} phases that are formed upon intercalation.

DOI: 10.1103/PhysRevB.66.035107 PACS number(s): 73.61.Wp, 71.20.Tx, 79.60.–i, 78.70.Dm

I. INTRODUCTION

After the extensive studies in the past several years of novel carbon nanostructures such as fullerenes or carbon nanotubes, many fundamental investigations are now carried out on the electronic properties of endohedral compounds. Particularly, charge transfer in metallofullerenes where the fullerene molecule can encage different species such as La, Y, Tm, Gd, or Sc has been widely investigated, as these materials are expected to display remarkable electronic and structural properties associated with this charge transfer from the metal to the carbon cage. More recently, the encapsulation of fullerene molecules into nanotubes, giving so-called peapods, and even the encapsulation of metallofullerenes into single wall nanotubes has been reported and should extend this interest since these novel materials might display extremely interesting electronic properties. Generally, for the metallofullerenes, an ionic interaction between the endohedral molecule and the carbon cage is to be seen as a crude picture and hybridization effects have to be taken into account.

Recently, the discovery was made that the introduction of a small amount of nitrogen into the metallofullerene reactor favors the formation of a new family of stable endohedral fullerenes \ce{A_xB_{3-x}N@C_{80}} \(x=0-3\). One prominent member of this new family is \ce{Sc_3N@C_{80}}. The nuclear magnetic resonance (NMR) spectrum of \ce{Sc_3N@C_{80}} has only two lines, consistent with \textit{I}\textsubscript{h} symmetry for the fullerene cage. It is well known that the symmetry of stable pristine \ce{C_{80}} is \textit{D}_{5\text{h}} (Ref. 12) and this is supported by calculations. These calculations also discuss the stability of the seven \ce{C_{80}} isomers allowed by the isolated pentagon rule, as a function of a charge transfer to the fullerene cage. On going from the neutral to a \textit{C}_{4\text{h}} state, the icosahedral \ce{C_{80}} is predicted to switch from being the least stable to the most stable carbon cage. This \textit{I}\textsubscript{h} structure could become even more energetically favorable in the \ce{C_{80}^-} state. Therefore it seems natural to suggest that the \ce{C_{80}} cage with \textit{I}\textsubscript{h} symmetry is stabilized by charge transfer from the nitride cluster in \ce{Sc_3N@C_{80}}. The exact amount of charge transfer, however, has up to now been unknown and should be determined experimentally, not only to settle this specific question, but also to further our knowledge of these novel encapsulated transition metal nitride fullerene clusters in general.
orbits. Furthermore, upon K intercalation several $K_i$(Sc$_3$N@C$_{80}$) phases are formed, as can be determined from the K 2$p$ photoemission doublet whose binding energy depends strongly on the chemical environment. The various metallofullerene salts are compared to both $K_i$(C$_{60}$) and $K_i$(C$_{70}$), regarding both their electronic structure and charge transfer characteristics.

II. EXPERIMENT

The preparation and separation of Sc$_3$N@C$_{80}$ is described elsewhere. Thin films of Sc$_3$N@C$_{80}$ were prepared for photoemission by sublimation onto a clean single crystalline Au(110) surface. The samples were then transferred under ultrahigh vacuum (UHV) conditions into a Perkin Elmer spectrometer, where they were studied using monochromatic Al $K\alpha$ radiation (1486.6 eV) with an energy resolution of 350 meV. The UPS was carried out using the He $L\alpha$ line (21.2 eV) from a helium discharge lamp with an overall energy resolution of 150 meV. X-ray absorption at the Sc $L_{2,3}$ edge was performed at the VLS-PGM beamline at BESSY; the resolution was 150 meV and the photon energy was calibrated from the binding energy (BE) of Au core level photoemission signals from the clean substrate.

Theoretical simulations of the Sc $L_{2,3}$ XAS spectra were performed using an atomic multiplet approach, which takes the presence of the core hole in the photoemission final state explicitly into account. Both the crystal field multiplet (CFM) interactions, as well as the covalency in the form of charge transfer (CT) have been taken into account. The CFM model is a single configuration calculation for Sc ions. The atomic interactions are included for a Sc$^{n+}$ ion and the chemical surroundings are simulated using an electrostatic crystal field. This method has been applied to Sc$^{3+}$ ions with a $3d^0$ configuration in simulations of Sc$_2$O$_3$ and ScF$_3$. For a $3d^1$ configuration Ti$^{3+}$ ions—which are isoelectronic with Sc$^{3+}$—have been calculated. The CT model extends the CFM model by using multiple configurations, e.g., $3d^1 + 3d^0 L$, where $L$ stands for an electron transfer from the surroundings (the “ligands,” L) to the central atom. The simplest way to arrive at the effective $3d$ count is by mixing two configurations in the calculations. This approach will be used here.

Intercalation was performed in UHV by evaporating potassium for 10 min from degassed commercial SAES getters. During intercalation the sample was held at 450 K. To quantify the degree of intercalation, we used the relative intensity of the K 2$p$/C 1$s$ photoemission peaks, taking into account the appropriate photoemission cross section of these states.

III. RESULTS AND DISCUSSION

A. Pristine material

1. The Sc valence state

The Sc$_3$N unit in Sc$_3$N@C$_{80}$ possesses 12 valence electrons in total. In the context of such an endohedral compound it is naturally of interest to ask how these charges are distributed between the constituents of the Sc$_3$N unit and the carbon cage.

Photoemission from the Sc 2$p$ levels can give a first rough estimation of the Sc valency from the chemical shift. The observed BE positions in Sc$_2$@C$_{84}$ and Sc$_3$N@C$_{80}$ are quite similar (see the inset to Fig. 1), which is the first hint that the Sc ions are close to a formally trivalent Sc (III) state. However, as the net BE of core levels depends not only on the degree of ionization of the atom concerned, but also on the screening of the core hole in the photoemission final state, we have performed XAS measurements at the Sc $L_{2,3}$ edge in order to try and determine the exact degree of charge transfer. As mentioned previously, XAS at the Sc $L_{2,3}$ edge has already been successfully used in this manner to determine the effective Sc valency in Sc$_2$@C$_{84}$ to be 2.6 (i.e., Sc$^{3d^{0.4}}$).

Figure 1 displays the XAS spectra of both Sc$_3$N@C$_{80}$ and Sc$_2$@C$_{84}$. The Sc$_3$N@C$_{80}$ spectrum shows a more intense Sc $L_3$ peak and new shoulders appear on the high energy sides of both main Sc peaks compared to Sc$_2$@C$_{84}$. These subtle differences show that the effective valency in both compounds is not exactly the same. This is related to a different interaction of the encaged cluster with the carbon cage and within the cluster itself. In order to quantify the effective valency, we compare the data with CFM-CT simulations, such as those shown in Fig. 2 for the Sc $L_{2,3}$ edges. The topmost spectrum displays the simulation for ionic Sc$^{2+}$ where the ground state is $[\text{Ar}]3d^1$, the bottommost spectrum the one of ionic Sc$^{3+}$ with the $[\text{Ar}]3d^0$ ground state. The experimental spectrum obviously does not match either of these extreme configurations but rather lies in between. This indicates that hybridization effects have to be taken into account. The best agreement between the simulation and the experimental data is achieved for a 60% admixture of the $3d^1$ configuration (Fig. 2), leading to an effective degree of ionization of close to 2.4 electrons per Sc, i.e., a formal
seven IPR isomers not only in the \((C_{80})^6\) charge state\(^{2,3,7,13-15}\) but also (to a lesser extent) in a \((C_{80})^4\) charge state.\(^{13}\)

The problem with the simple picture given in the opening paragraph of this section is that the experimental XAS data offer no support for a simple \(3d^{0.6}Sc^{3+}\)-like scenario: there is clearly finite \(3d\) electron density at the Sc site. The key here is to take the short Sc–N bond length into account (1.98 Å in the endohedral vs 2.25 Å in bulk ScN—Ref. 19). As ScN is quite an ionic compound, the shorter Sc–N distance in the endofullerene is a signal of significant Sc/N covalence. This sharing of electron density will naturally lead to increased charge density at the Sc, thus accounting for the \(3d^{0.6}\) configuration of the Sc ions. This hypothesis receives support from the planar (i.e., not ammonia-like) form of the Sc\(_3\)N cluster, which is consistent with a significant \(p_{\pi}-d_{\pi}\) ("back-donation") component in the Sc–N bonding,\(^{18}\) in which only 0.3 electrons per Sc are fully transferred to N.\(^{20}\) As a result of the observed formal \(3d^{0.6}\) configuration in Sc\(_3\)N@C\(_{80}\), and based on the foregoing one can propose that \((3 \times 2.4) - (3 \times 0.3) = 6.3\) electrons are available for transfer from a Sc\(_3\)N cluster on its encapsulation in a C\(_{80}\) molecule,\(^{21}\) thus providing the six electrons required to optimally stabilize the \(I_h\) C\(_{80}\) cage as discussed previously.

In an alternative view of the Sc–N bonding one electron per Sc would be considered to be fully transferred to N (and 0.6 \(3d\) electrons per Sc remain on the transition metal). Then \((3 \times 2.4) - 3 = 4.2\) electrons are available for transfer to the C cage. Considering the short Sc–N distance in this system, however, it is likely that in this way the N electron density is underestimated, as some of the Sc \(3d^{0.6}\) electron density is certainly participating in the \(p_{\pi}-d_{\pi}\) covalent bonding. Consequently, the N would then have, in fact, more than the eight electrons it requires in its outer shell. However, pursuing this avenue further for the sake of argument, one could thereby arrive at a net stabilization of the \(I_h\) C\(_{80}\) isomer in its four minus charge state,\(^{13}\) which would be needed to explain the \(^{13}C\) NMR data.\(^{10}\) Incidentally, a four minus charge on the host cage would also be one way to explain the different retention time of Sc\(_3\)N@C\(_{80}\) in chromatographic separation compared to systems such as La\(_3\)@C\(_{80}\), which are taken to be close to the ionic limit of sixfold electron transfer\(^{10}\) (although the results of Refs. 22 and 1 warn us that an ionic picture even for La@C\(_{82}\) is an oversimplification). Nevertheless, both the issues regarding the undercounting of the effective electron density accessible to the nitrogen given previously and the large binding energy of the highest occupied electronic states of solid Sc\(_3\)N@C\(_{80}\) observed in the valence band photoemission data presented in the following are strong arguments against this “four minus” picture and in favor of a \((C_{80})^5\) scenario.

### 3. Valence band photoemission

In valence band photoemission spectroscopy the matrix-element weighted occupied density of states is probed, and one can thereby gain information not only regarding the energy distribution of the electronic states but also about the symmetry and the stability of the metallofullerene in ques-
The He I photoemission spectra shown in Fig. 3 are mainly due to emission from the carbon cage states, a conclusion drawn from photoionization cross-section considerations. For Sc$_3$N@C$_{80}$ the ratio between atomic photoionization cross sections for C 2p, Sc 3d, and N 2p emission leads to an upper limit for the contribution from the Sc$_3$N unit to the overall photoemission response upon using He I radiation of only about 2%. Generally, the valence bands of fullerences and metallofullerenes display broad structures in photoemission, this breadth reflecting the splitting of electronic states due to the (generally) low symmetry of the carbon cage and due to covalent interactions between the encaged metal and the host molecule. As typical examples, the spectra of Sc$_2$@C$_{84}$ and C$_{84}$ are also depicted in Fig. 3. In contrast, Sc$_3$N@C$_{80}$ exhibits four narrow peaks in the low energy region (BE < 5 eV) of its valence band photoemission spectrum. This would be fully consistent with the suggestion that the carbon cage possesses high, icosahedral symmetry. Indeed, the spectrum of Sc$_3$N@C$_{80}$ is more comparable to that of C$_{60}$ ($I_h$) than to that of pristine C$_{80}$. In conclusion, compared to pristine C$_{80}$ which has low, $D_2$ symmetry, we observe well-defined, sharp photoemission features for Sc$_3$N@C$_{80}$ pointing to the high degeneracy of the electronic states, concomitant with $I_h$ symmetry of the fullerene cage. The fact that the Sc$_3$N@C$_{80}$ molecule as a whole has only $C_3$ symmetry (or even is reduced to $C_1$ in the solid state) evidently does not give rise to a lifting of the degeneracy of C$_{80}$’s molecular orbitals visible above the vibronic broadening which dominates the width of the individual molecular orbitals in valence band photoemission. Thus, in the Sc$_3$N@C$_{80}$ molecule, the majority of the electronic levels remain quite tightly bunched in energy.

A further point of interest in the valence band spectrum of Sc$_3$N@C$_{80}$ is the fact that the onset of the HOMO is at the relatively large binding energy of $\sim 1$ eV. The well-known electron correlation effects in the fullerenes$^{23,24}$ mean that in an ionizing spectroscopy such as photoemission this onset energy cannot necessarily be interpreted simply as an expression of the minimal energetic separation of the HOMO and LUMO. Nevertheless, the BE of this onset is too large to be consistent with an odd-electron system, such as is approximated in certain endohedrals such as M@C$_{82}$ [M=Gd,La]. The onset energy is also such that it presents a strong argument in favor of a (C$_{80}$)$^{6-}$ scenario in Sc$_3$N@C$_{80}$, as the further Jahn–Teller splitting of a threefold degenerate LUMO (such as that of the C$_{80}$ molecule) containing four electrons would not be able to account for the magnitude of the observed gap in the electronic excitation spectrum, whereas a sixfold electron transfer to the $I_h$ C$_{80}$ cage would result in a closed-shell structure with the excitation gap then corresponding to a large extent to the (large) LUMO − LUMO + 1 energy separation. We note that the theoretical predictions of an energy gap of about 2 eV for both (C$_{80}$)$^{6-}$ and Sc$_3$N@C$_{80}$ are in qualitative agreement with our observation.$^{15,18}$

### B. K-doped material

This part of the paper deals with additional (so-called “combinational”) doping of the metallofullerene Sc$_3$N@C$_{80}$ by K intercalation. From x-ray photoemission (XPS) measurements, the intensity ratio between the K 2p and C 1s lines can give a good indication of the overall potassium concentration in such intercalation compounds. Figure 4 displays the K 2p photoemission lines for several intercalation levels in K$_x$(Sc$_3$N@C$_{80}$). Similar to pristine fullerenes, many metallofullerenes crystallize at room temperature in a fcc-like close-packed structure.$^{25}$ Hence, both octahedral and tetrahedral interstitial sites can serve as hosts for the K ions (respectively, labeled O and T in Fig. 4) and this leads to different K 2p peak positions in the photoemission spectra due to their different Madelung potentials.$^{26,27}$ In previous
data from K intercalated fullerenes, the features evolving upon intercalation have been related to the phase diagrams of K$_x$C$_{60}$ and K$_x$C$_{70}$. Thus, in analogy with the situation in these systems, the observation of two K 2$p$ doublets—indicating the occupation of two different interstitial lattice sites—right from the onset of intercalation, is consistent with an fcc-like structure for Sc$_3$N@C$_{80}$. The doublets corresponding to the octahedral and tetrahedral sites are observed at 298 and 295.3 eV, and at 296.9 and 294.1 eV, respectively (with the energy separation of each component of the same doublet equal to the well known spin–orbit splitting of the K 3$p_{1/2}$ and K 3$p_{3/2}$ levels). For an average stoichiometry K$_{0.7}$ (Sc$_3$N@C$_{80}$), both tetrahedral and octahedral sites are occupied but the latter site is favored (see the bottommost spectrum of Fig. 4). In K$_x$C$_{60}$ for x<3, at room temperature, phase separation between C$_{60}$ and K$_x$C$_{60}$ is observed, giving rise to a constant O to T intensity ratio of 1:2, independent of the average potassium stoichiometry. Only above 450 K is a stable phase K$_x$C$_{60}$ observed, in which only the octahedral site is occupied. A similar kind of phase separation is observed for K$_x$C$_{70}$ and K$_x$C$_{70}$ phases. At room temperature, stable K$_x$C$_{70}$ phases are observed for x = 1, 4 and 6. In the light of this behavior in the system K$_x$C$_{70}$, the predominant occupancy of octahedral site in K$_{0.7}$ (Sc$_3$N@C$_{80}$), signaled by the more intense O component in Fig. 4, is probably correlated to the formation of a K$_x$(Sc$_3$N@C$_{80}$) phase.

Upon further intercalation, the O site feature reduces in intensity with respect to the T peak, consistent with the formation of a fcc-like K$_x$(Sc$_3$N@C$_{80}$) phase. This behavior is in contrast to doped C$_{70}$ where a switch from a fcc- to a bcc-like lattice is observed for x>1. In our case, only for higher doping x>3, does such a transformation into a body centered structure appear to occur. This is similar to the K$_x$C$_{60}$ case and is confirmed by the observation of a new K 2$p$ doublet (marked as $T_2$ at 297.2 and 294.3 eV in Fig. 4), related to a body centered crystal symmetry. This is consistent with the formation of a stable phase K$_x$(Sc$_3$N@C$_{80}$) or K$_x$(Sc$_3$N@C$_{80}$) as in K$_x$C$_{60}$. This observed doping dependence is also typical of other metallofullerenes, as can be seen by the example of potassium intercalated Tm@C$_{82}$. Additional information can be extracted from the C 1$s$ core level of the C$_{80}$ cage and the N 1$s$ and Sc 2$p$ core level spectra of the nitride unit. We deal first with the C 1$s$ spectrum of the pristine endohedral before going into the doping dependence. C$_{80}$, with its icosahedral molecular symmetry, possesses only a single C site, giving rise to a C 1$s$ core level photoemission line shape which is symmetric with a full width at half maximum (FWHM) of 0.65 eV. In the preceding, strong arguments have been presented—over and above the $^{13}$C NMR data—for the icosahedral molecular symmetry of the C$_{80}$ cage in Sc$_3$N@C$_{80}$. Empty I$_h$ C$_{80}$ possesses two inequivalent C sites, whereas the endohedral molecule has lower overall symmetry due to the presence of the encapsulated Sc$_3$N unit. The latter has indeed been suggested to interact in a bonding sense via the Sc atoms with the corannulene units of the carbon cage, but nevertheless does not lead to a significant distortion of the C$_{80}$ cage itself. This cluster–cage interaction could well be the source of the larger FWHM of the C 1$s$ core level line of 1.2 eV for the C$_{80}$ endohedral, compared to the FWHM of 0.7 eV recorded for C$_{70}$. An empty fullerene also possessing two inequivalent C sites. However, the results of the valence band photoemission, taken together with the absence of a strong distortion of the C$_{80}$ host cage suggests that the cluster–fullerene interaction does not lead to a large energetic splitting between the molecular orbital-derived electronic states which would be formally degenerate in a hypothetical I$_h$ Sc$_3$N@C$_{80}$ molecule.

Returning to the question of the doping dependence of the core level spectra, we note that the C 1$s$ peak evolution in Sc$_3$N@C$_{80}$ upon intercalation, displayed in Fig. 5, is completely different compared to the situation found in C$_{60}$ and C$_{70}$, with the C 1$s$ peak sharpening upon intercalation (the width being reduced by about 8% on going from x = 0 to x = 3.1). Therefore, in contrast to the C$_{60}$ intercalation compounds, doping possibly leads to a higher degree of equivalence for the carbon sites of the C$_{80}$ cage. We also note that the peak remains nearly symmetric at every stage of intercalation, unlike in the case of K–C$_{70}$ and unlike K–C$_{60}$.

Furthermore, unlike both intercalated C$_{60}$ and C$_{70}$, the peak position shifts toward higher binding energy for x = 2.1 (from 285.2 to 285.3 eV) and then goes back to 285.25 eV for x = 3.1. The origin of such a nonrigid shift could be related to the pinning of the Fermi level in the different K$_x$(Sc$_3$N@C$_{80}$) phases. In addition, the individual separation between the newly occupied HOMO levels and the Fermi level is strongly altered by the local C$_{80}$ cage structure, with the consequence that the shift of the C 1$s$ line cannot be explained within a simple rigid band model. A more accurate analysis is required with the aid of sample distillation, for instance, in order to get more precise information on the phases that are present in the crystal. Until this is carried out, the behavior of the C 1$s$ BE upon doping remains an open question.
We now discuss the changes of the core level data from the encaged cluster upon intercalation. Figure 6 shows the Sc 2p and N 1s peaks as a function of K content. It is obvious from the data that these electronic states are scarcely altered by the doping process. For increasing potassium concentration, a small broadening of the core level spectra of about 250 meV is observed. This broadening can be explained by screening effects due to the polarizable potassium counterions: similar effects have been seen in Tm 4d and Tm 4f photoemission from KxTm@C82.30 For the Sc 3d absorption edge in XAS, no changes of the line shape and of the peak intensities are observed upon intercalation. This makes it clear that the effective Sc valency is not altered with intercalation-induced doping of the metallofullerene. This is in good agreement with recent doping experiments on other metallofullerenes such as Sc2@C84, Tm@C82, Gd@C82, Ce2@C74.30

The valence band photoemission spectra at various K intercalation levels are presented in Fig. 7. At very low doping, the Fermi level can be assumed to be pinned near the edge of the lowest unoccupied molecular orbital (LUMO),31 thus making the onset observed at 1.1 eV a more accurate estimation of the band gap relevant for ionization of the molecule. From x = 0.7 and upwards, filling of the LUMO gives rise to a broad structure at about 1.5 eV binding energy. Subsequent doping leads to the appearance of several Kx(Sc3N@C80) phases, with the superposition of their different photoemission signatures giving rise to broad spectral structures. Thus the very sharp peaks observed near the Fermi level in the pristine material merge into structureless features in the valence band of the intercalated material.

IV. CONCLUSIONS

We have presented a systematic study of the electronic structure and charge transfer characteristics of the novel trinitride endohedral fullerene Sc3N@C80 in the solid state, both in its pristine and K intercalated forms.

As regards the pristine endofullerene, existing spectroscopic and theoretical results,18 taken together with the Sc L2,3 XAS (experiment and cluster-based simulation) and photoemission data presented here show the following:

1. the effective 3d electron count at the Sc sites is 0.6 electrons (compared to a Sc 3d0.4 configuration in Sc2@C84),
2. there is strong covalent bonding between the Sc and the N in the encapsulated cluster,
3. the onset in photoemission of the highest occupied molecular orbitals is ~1 eV below EF,
4. the valence band photoemission spectrum of solid Sc3N@C80 shows a wealth of fine structure, more reminiscent of that of C60 than of a typical higher-fullerene endohedral,
5. the C 1s core level spectrum of Sc3N@C80 shows a relatively narrow main line for an endohedral fullerene, whose fine-structure suggests the presence of only two inequivalent C sites.

These facts can be consistently explained in a picture in which a significant degree of dπ−pπ covalent “backbonding” within the Sc3N unit is responsible for the finite 3d electron density at the Sc site, with the remaining valence electron density from the Sc3N cluster being transferred to fully populate the lowest lying C2π*-derived pπ* molecular orbital. This leads, in terms of C80’s LUMO population with respect to the neutral molecule, to a hexaanionic state for the fullerene cage. This (C80)6− state is the clue to understanding the photoemission data, as both the narrow C 1s line as well as the large gap in the valence band photoelectron spectrum are to be expected only for the case in which the struc-
ture of the C$_{80}$ host cage has high, $I_h$ symmetry and a fully closed shell. This C$_{80}$ isomer is highly unstable in its neutral form, but is predicted to be the most stable isomer in a hexaannionic state. Thus, the data presented here, together with the solution $^{13}$C NMR results$^{10}$ serve to put the $I_h$ identity of the C$_{80}$ carbon cage in this endohedral beyond doubt.

In addition, by analyzing the evolution of the K 2p core level photoemission features as a function of intercalation, and comparison of the results with the K intercalated C$_{60}$ and C$_{70}$ systems, we were able to give evidence compatible with the formation of several distinct phases in the K$_x$(Sc$_3$N@C$_{80}$) system.

**ACKNOWLEDGMENTS**

This work was supported by the EU-sponsored TMR Research Network “FULPROP” (ERBFMRXCT-970155). In addition, T.S. thanks the BMBF (05 5F8BD11) and T.P. the ÖAW and FWF for funding. Our thanks also to Gotthardt Seifert for useful discussions.

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