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Metal–ring interactions in actinide sandwich compounds: A combined normalized elimination of the small component and local vibrational mode study

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ABSTRACT
The relativistic Hamiltonian Normalised Elimination of the Small Component with atomic unitary transformation (NESCau) was used for the first time to calculate geometries and harmonic vibrational frequencies of actinide sandwich compounds \( \text{An}(C_nH_n)_2 \) (An: Th, Pa, U, Np, Pu; \( n = 5–8 \)). In addition, the Local Vibrational Mode analysis, the Natural Bond Orbital analysis, and the Atoms in Molecules analysis were applied to quantitatively assess the intrinsic strength and the nature of the An-ring interactions. Our results show that actinide sandwich compounds prefer strong covalent interactions between actinide and carbon atoms, similar to those found for homologous ferrocene-type complexes \( \text{Fe}(C_nH_n)_2 \). The metal centre rather than the ring size plays a key role in the strength of those interactions.

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NESCau; actinides; sandwich compounds; local mode analysis; intrinsic bond strength

1. Introduction
The discovery of ferrocene, \( \text{Fe}(C_5H_5)_2 \), in 1951 [1], added a new important field to organometallic chemistry: metal sandwich complexes [2]. The first synthesis of sandwich complexes using an actinide metal centre was reported in 1968 [3] followed by the series \( \text{M}(C_8H_8)_2(M = \text{Th}, \text{U}, \text{Np}, \text{Pu}) \) [4,5]. Organometallic complexes of actinide-\( \eta^3 \)-carbocyclic derivatives were added to the repertoire because they showed promising magnetic, electronic, photo-physical and catalytic properties [6–9]. The last few years have seen a significant increase of the synthesis of thorium and uranium sandwich complexes [10–14]. In addition, neptunium and plutonium complexes with four \( (C_7H_7) \) and two \( (C_8H_8) \) ligands have shown a unique behaviour in many small-molecule reactions [15,16]. In order to further develop the field of actinide sandwich complexes, a deeper and more comprehensive understanding of f-block metal-ligand bonding is needed. The experimental investigation of actinide compounds is extremely challenging because of their harmful radioactive character. Also computational studies are difficult, because they require a careful...

consideration of relativistic effects, which play a significant role for the modelling and prediction of reliable physical and chemical properties of compounds with heavy elements, i.e. atoms with an atomic number larger than 20 \([17,18]\), as pointed out in the pioneering work of Desclaux \([19]\), Grant \([20]\) and Pyykkö \([21,22]\). One has also to consider that relativistic calculations may significantly increase the computational costs; therefore it is imperative to find the right balance between accuracy and efficiency.

Over the past decades a lot of effort has been put into the development of quantum chemical methods including relativistic effects \([23–41]\). In analogy with Perdew’s prominent ‘Jacob’s ladder’ of density functional theory \([42]\) one can establish a relativistic Jacob’s ladder, as sketched in Figure 1 stretching from the non-relativistic earth to the quantum electrodynamics (QED) heaven \([43,44]\).

The most rigorous way to include relativity in the calculation of molecular systems is to use Dirac’s full 4-component (4c) formalism leading to wave functions being vectors of four complex numbers (known as bispinors) \([45,46]\). A large variety of approximate methods (the lower ladders in Figure 1) have been derived over the years, motivated by the assumption that the full 4c approach is computationally too demanding and cannot be applied to larger molecules. One of the most popular approximation is the so-called 2-component (2c) approximation, derived from decoupling the large and small components of the Dirac spinor. This approach forms the basis for two important families; the Douglas-Kroll-Hess (DKH) \([47,48]\) method and the regular approximation (RA) \([49]\). The most famous DKH-type Hamiltonian is the second-order DKH (DKH2) \([47,48]\), which encouraged the development of higher finite-order Hamiltonians, the so-called DKHn Hamiltonians \([50–52]\). Concerning the RA family, the zeroth-order RA (ZORA) has been widely used in different forms \([53,54]\). It is important to note that these low-order approximate relativistic Hamiltonians do not treat the core orbitals exactly.

In 1997, Dyall developed the Normalised Elimination of the Small Component (NESC) method \([23,24]\), an important milestone, as NESC could be identified as the first Dirac-exact 2-component relativistic approach (X2C), i.e. a quasi-relativistic (2c) method, which reproduces the one-electron energies of the original 4-component Dirac method and treats both core and valence orbitals exactly \([55]\). Dyall’s work was the starting point for the development of a whole family of X2C relativistic methods \([56–61]\), also called Dirac-exact NESC methods \([62–69]\), defining the upper ladders in Figure 1. Nowadays, X2C methods are the most popular exact relativistic methods for the characterisation of compounds with heavy atoms \([26]\). Another well-known exact relativistic method is the infinite-order DKH (IODKH; also called infinite-order two-component) method \([70,71]\), which is based on the Barysz-Sadlej-Snijders (BSS) approach \([72]\). A comprehensive review of those approaches are collected in Ref. \([73]\).

Since the X2C Hamiltonian is mathematically simpler, analytic nuclear gradients \([27–29,74]\) and analytic Hessians \([31,32,56]\) have been derived, being important for relativistic geometry optimisations and frequency calculations. Recently, we adopted a new version of NESC using the atomic approximation (NESCau), which substantially increases the efficiency of relativistic calculations \([56]\), as demonstrated in this work.
In addition to choosing the appropriate relativistic method for the description of the actinide sandwich compounds, a suitable measure assessing the intrinsic strength of the An-ring interaction is needed. The local vibrational mode analysis, originally developed by Konkoli and Cremer [75–78] provides the perfect tool for this purpose. So far, the local mode analysis has been successfully applied for the characterisation of covalent bonds [79–85], a broad range of weak chemical interactions in gas phase, solution and an enzyme environment [86–102], as well as chemical bonding in periodic systems and crystals [103,104]. In this work, we extended the local vibrational mode analysis by introducing a new quantitative measure of the intrinsic strength of metal–ring interactions in metal sandwich compounds, as is further described below.

A test set of 20 An(C_nH_n)\_2 (n = 5–8; An = Th, Pa, U, Np, Pu) complexes (see Figure 2) was investigated in this work with NESCau followed by the local mode analysis, in order to deepen our understanding of actinide–ligand bonding, focusing in particular on the following objectives:

- to determine the strength of the An-ring interaction in An(C_nH_n)\_2 utilising the local mode analysis;
- to investigate how the strength of the An-ring interaction is related to 5f orbital occupation;
- to determine the intrinsic strength of the An-C ring bonds, in particular their covalent or non-covalent character;
- to compare metal–ring interactions of An(C_nH_n)\_2 with those of the Fe(C_3H_5)\_2 prototype.

The paper is arranged as follows. In Section 2, we provide details about the computational methods applied including a description of NESCau and the local mode analysis. In Section 3, we systematically elucidate the An-ring interactions and discuss the most important factors determining the stability of these complexes; including local stretching force constants as bond strength descriptors, the natural bond orbital (NBO) analysis [105] exploring 5f occupation, and the topological analysis of the electron density analysing the covalent character of An-ring bonding. The last section summarises the most important results and draws some conclusion which are useful for future fine-tuning and design of actinide sandwich compounds.

2. Computational method

2.1. Normalised elimination of the small component

The continuous development of NESC has been described in several articles [29,32,62–65,67,106,107] including two reviews [26,66], and a number of applications have been reported [27,68,69,108,109]. Therefore, in the following only a short introduction of NESC is given focusing on the NESCau extension.

Starting from the 4c-Dirac equation with embedded restricted kinetic balance (RKB), a Dirac equation in matrix picture was obtained by Dyall [23],

\[
\begin{pmatrix}
V & T \\
T & W - T
\end{pmatrix}
\begin{pmatrix}
A_- \\
B_-
\end{pmatrix}
= \begin{pmatrix}
S & 0 \\
0 & \frac{1}{2c^2}T
\end{pmatrix}
\begin{pmatrix}
A_+ \\
B_+
\end{pmatrix}
\begin{pmatrix}
\varepsilon_- & 0 \\
0 & \varepsilon_+
\end{pmatrix}
\]

(1)

where \(S, T\) and \(V\) are, respectively, the overlap, kinetic energy and nuclear attraction integral matrices, \(W\) is the matrix of the operator \((\sigma \cdot p)V(r)(\sigma \cdot p)/(4c^2)\), and \(\varepsilon_+\) (\(\varepsilon_-\)) is the electronic (positronic) eigenvalues with the eigenvectors \(A_+\) (\(A_-\)) for large component and \(B_+\) (\(B_-\)) for pseudo-large component. Chemists are generally not interested in the positronic solutions in \(\varepsilon_-\), i.e., they are discarded. After some algebra, Dyall derived the NESC equation considering only the electronic eigenvalues \(\varepsilon_+\):

\[
\tilde{L} A_+ = \tilde{S} A_+ \varepsilon_+
\]

(2)

with the NESC Hamiltonian

\[
\tilde{L} = U^\dagger T + TU - U^\dagger(T - W)U + V
\]

(3)
and the metric
\[ \tilde{S} = S + \frac{1}{2c^2} U^T T U. \] (4)

In above equations, \( U \) can be calculated by \( B_+ = UA_+ \).

Within the one-electron approximation to the many-body relativistic problem, the NESC Hamiltonian \( \tilde{L} \) has to be renormalised on the non-relativistic metric \( S \) through the following equation
\[ H_{\text{NESC}} = G^\dagger \tilde{L} G \] (5)
where the renormalisation matrix [60] is given by
\[ G = S^{-1/2} K S^{1/2}, \] (6)
\[ K = \left( S^{-1/2} \tilde{S} S^{-1/2} \right)^{-1/2}. \] (7)

The renormalised NESC Hamiltonian \( H_{\text{NESC}} \) in Equation (5) is then used to replace the one-electron Hamiltonian in the non-relativistic Fock matrix leading to the coverage of relativistic effects. This is the essence of the Dirac-exact NESC method [106], also called X2C method, as stated [110].

Compared to other relativistic methods, NESC (i.e. X2C) has the advantage of accuracy, simplicity and efficiency [59]. However, for medium-sized and larger molecules, NESC becomes time-consuming due to the involved matrix inversion and diagonalisation procedures. An efficient cure for this problem is the so-called From Atoms To Molecule approach [59]. Considering that the molecular \( U \) matrix is substantially atomically local, it can be approximated by the direct sum of a series of atomic \( U \) matrices; similarly the molecular \( G \) matrix can also be constructed approximately. This NESC variant, which is essentially as accurate as standard NESC, has been coined NESC with atomic unitary transformation (NESCau, i.e. X2Cau) [56]. Since the approximate \( U \) and \( G \) matrices in NESCau do not depend on molecular geometries, analytic calculations of nuclear gradients and Hessians are very efficient [56], and therefore can be used to investigate large molecules as is demonstrated in this study.

### 2.2. The local vibrational mode analysis

The central starting point is Wilson's theory of vibrational spectroscopy. In Equation (8), the Wilson equation of vibrational spectroscopy for a vibrating systems with \( N \) atoms is given [111–117]:
\[ F^\dagger \tilde{L} = M LA \] (8)
where \( F^\dagger \) is the force constant matrix or Hessian matrix expressed in Cartesian coordinates \( x_i \) (\( i = 1, \ldots, 3N \)). \( M \) is the mass matrix and matrix \( \tilde{L} \) collects the vibrational eigenvectors \( \lambda_\mu \) in its columns (\( \mu = 1, \ldots, N_{\text{vib}} \)). The number of vibrational modes \( N_{\text{vib}} \) equals \( 3N - N_{\text{tr}} \) with the translational and rotational modes \( N_{\text{tr}} \) being 5 for linear and 6 for non-linear molecules. \( A \) is a diagonal matrix with the eigenvalues \( \lambda_\mu \), which leads to the (harmonic) vibrational frequencies \( \omega_\mu \) according to \( \lambda_\mu = 4\pi^2 c^2 \omega_\mu^2 \). The tilde symbol indicates mass weighting. The normal mode eigenvectors and eigenvalues are obtained by diagonalising the force constant matrix \( F^\dagger \) defined in Equation (8) according to \( \tilde{L}^\dagger F^\dagger \tilde{L} = \Lambda \) and \( \tilde{L}^\dagger M \tilde{L} = I \).

One usually re-normalises the normal mode vectors \( \tilde{L}_\mu \) according to \( L = \tilde{L} (M^R)^{1/2} \), where the elements of the mass matrix \( M^R \) are given by \( m^R_\mu = (\tilde{L}_\mu^\dagger \tilde{L}_\mu)^{-1} \) and represent the reduced mass of mode \( \mu \).

Without mass-weighting Equation (8) takes the form [111,112,116]
\[ F^\dagger L = MLA \] (9)
leading to the diagonal normal force constant matrix \( K \), and the reduced mass matrix \( M^R \) in normal coordinates \( Q \), respectively.
\[ L^\dagger F^\dagger L = K \] (10)
\[ L^\dagger M L = M^R \] (11)

The dimension of matrices \( K \) and \( M^R \) is \( N_{\text{vib}} \times N_{\text{vib}} \).

It is often more convenient to express the molecular geometry in terms of internal coordinates \( \mathbf{q} \) rather than Cartesian coordinates \( \mathbf{x} \). Doing so the Wilson equation adopts the following form [111]:
\[ F^\dagger D = G^{-1} DA \] (12)
where \( D \) collects the normal mode vectors \( d_\mu \) (\( \mu = 1, \ldots, N_{\text{vib}} \)) column-wise, and the Wilson matrix \( G \), which is defined as
\[ G = BM^{-1}B^\dagger \] (13)
represents the kinetic energy in terms of internal coordinates. The elements of the \( B \) matrix in Equation (13) are defined by the partial derivatives of internal coordinates with regard to Cartesian coordinates. It is important to note that the \( B \) matrix plays a central role for the Wilson equation of spectroscopy, namely connecting different sets of coordinates (internal, symmetry, curvilinear, etc.) with the Cartesian coordinates [111]. Therefore, whenever a new set of coordinates is introduced, the first step is to derive the appropriate \( B \) matrix. This is an important point for deriving a new bond strength measure for...
metal–ring interactions, as discussed below. Diagonalisation of Equation (12) leads to

\[ \mathbf{D}^{\dagger} \mathbf{F}^{a} \mathbf{D} = \mathbf{K} \quad (14) \]

Equations (9) and (12) are connected by the following equations [111]:

\[ \mathbf{F}^{a} = \mathbf{C}^{\dagger} \mathbf{F}^{aC} \mathbf{C} \quad (15) \]
\[ \mathbf{D} = \mathbf{B} \mathbf{L} \quad (16) \]

Matrix \( \mathbf{C} \) is the pseudoinverse of the \( \mathbf{B} \) matrix defined by

\[ \mathbf{C} = \mathbf{W} \mathbf{B}^{\dagger} \left( \mathbf{B} \mathbf{W} \mathbf{B}^{\dagger} \right)^{-1} \quad (17) \]

where \( \mathbf{W} \) is an arbitrary nonsingular \( 3N \times 3N \) square matrix. At a stationary point (i.e., the energy gradient is a zero-vector), the \( N_{tr} \) translational and rotational eigenvectors are decoupled from the \( N_{vib} \) vibrational eigenvectors, and therefore \( \mathbf{W} \) does not affect the results [118]. For reasons of simplicity, one often uses \( \mathbf{W} = \mathbf{I}_{3N} \) (usually for the geometry optimisation procedure in internal coordinates) or \( \mathbf{W} = \mathbf{M}^{-1} \). Using the latter definition, which is more physically sound [119] this leads to

\[ \mathbf{C} = \mathbf{M}^{-1} \mathbf{B}^{\dagger} \mathbf{G}^{-1} \quad (18) \]

and

\[ \mathbf{B} \mathbf{C} = \mathbf{I}_{N_{vib}} \quad (19) \]

However, it should be noted that \( \mathbf{C} \mathbf{B} \neq \mathbf{I}_{3N} \) since \( \mathbf{B} \) is spanned in a \( N_{vib} \)-dimensional vibrational space.

While the solution of the Wilson equation, i.e. the transformation to normal coordinates \( \mathbf{Q} \) leads to a diagonal normal force constant matrix \( \mathbf{K} \) free from any electronic coupling, there is still mass-coupling which prevents the direct use of normal mode force constants as intrinsic bond strength measure, a fact which often has been overlooked. In 1998, Konkoli and Cremer [75,76,120–122] derived for the first time local vibrational modes directly from normal vibrational modes by solving the mass-decoupled Euler-Lagrange equations, i.e. by solving the local equivalent of the Wilson equation for vibrational spectroscopy. They developed the leading parameter principle [75,120] which states that for any internal, symmetry, curvilinear, etc. coordinate a local mode \( \mathbf{a}_{n} \) can be defined. \( \mathbf{a}_{n} \) is independent of all other internal coordinates used to describe the geometry of a molecule, which means that it is also independent of using redundant or non-redundant coordinate sets. The local mode vector \( \mathbf{a}_{n} \) associated with the \( n \)th internal coordinate \( q_{n} \) is defined as [75,120]

\[ \mathbf{a}_{n} = \frac{\mathbf{K}^{-1} \mathbf{d}_{n}^{\dagger}}{\mathbf{d}_{n} \mathbf{K}^{-1} \mathbf{d}_{n}^{\dagger}} \quad (20) \]

where the local mode \( \mathbf{a}_{n} \) is expressed in terms of normal coordinates \( \mathbf{Q} \) and \( \mathbf{d}_{n} \) is the \( n \)th row vector of the \( \mathbf{D} \) matrix defined in Equation (16).

\[ \mathbf{d}_{n} = \mathbf{b}_{n} \mathbf{L} \quad (21) \]

Equation (20) reveals that only matrices \( \mathbf{K} \) and \( \mathbf{D} \) are needed to determine \( \mathbf{a}_{n} \), i.e. once the normal analysis is completed, a following local mode analysis is straightforward [75,120].

To each local mode \( \mathbf{a}_{n} \) local mode properties can be assigned. The local mode force constant \( k_{n}^{a} \) of mode \( n \) (superscript \( a \) denotes an adiabatically relaxed, i.e. local mode) is obtained via Equation (22):

\[ k_{n}^{a} = \mathbf{a}_{n}^{\dagger} \mathbf{K} \mathbf{a}_{n} = (\mathbf{d}_{n} \mathbf{K}^{-1} \mathbf{d}_{n}^{\dagger})^{-1} \quad (22) \]

It is noteworthy that local mode force constants, contrary to normal mode force constants, have the advantage of being independent of the choice of the coordinates used to describe the molecule in question [75,76].

The local mode mass \( m_{n}^{a} \) of mode \( n \) is given by

\[ m_{n}^{a} = 1/G_{n,n} = (\mathbf{b}_{n} \mathbf{M}^{-1} \mathbf{b}_{n}^{\dagger})^{-1} \quad (23) \]

where \( G_{n,n} \) is the \( n \)th diagonal element of the Wilson \( \mathbf{G} \) matrix. For a chemical bond A–B, Equation (23) leads to \( M_{A}M_{B}/(M_{A} + M_{B}) \), which has the same form as the reduced mass of diatomic molecules.

Local mode force constant and mass are needed to determine the local mode frequency \( \omega_{n}^{a} \)

\[ (\omega_{n}^{a})^{2} = \frac{1}{4\pi^{2}c^{2}} k_{n}^{a} G_{nn} \quad (24) \]

Apart from these properties, it is straightforward to determine other local mode properties, such as infrared intensities [77].

In the following, we will derive a local mode stretching force constant for the An–ring interaction. One could think of calculating the local modes for all An–ring carbon interactions and averaging over the corresponding local mode force constants. A better way is to use only one interaction parameter describing the local stretching between the An atom and the centre M of the ring, as depicted in Figure 3 leading to a straightforward comparison of the strength of An–ring interaction in different An-sandwich complexes.

Similar to the stretching between two atoms [111], the \( \mathbf{B} \)-matrix of the stretching between the An atom and the
to examine the covalent character of the An–Cbonds 
in order to determine the 5f orbital occupation number of the An atom in each compound, in this way identifying which 5f orbital is participating in bonding. In order to examine the covalent character of the An–C bonds we applied the Cremer-Kraka Criterium for covalent bonding, which is based on the local energy density

\[ H(r) = G(r) + V(r) \]

where \( G(r) \) is kinetic energy density (positive, destabilising) and \( V(r) \) is potential energy density (negative, stabilising). Taken at the bond critical point \( r_b \) of the electron density \( \rho(r) \) between two bonded atoms, \( H(r_b) < 0 \) indicates a covalent bond, while \( H(r_b) > 0 \) indicates an electrostatic interaction.

The set of 20 complexes An\((C_nH_n)_2\) (An : Th(IV), Pa(V), U(IV), Np(IV), Pu(IV); \( C_nH_n = [C_5H_5]^-, [C_6H_6]^-, [C_7H_7]^+, [C_8H_8]^2- \)) and as a reference Fe\((C_5H_5)_2\) were calculated at the NESCau/M06 level of theory [132] utilising the Jorge-triple zeta valence quality plus polarisation (Jorge-TZP-DKH) basis set for all metals and the double zeta (Jorge-DZP-DKH) for carbon and hydrogen atoms [133–136]. These basis sets have been reported as an efficient choice for relativistic calculations of actinide compounds, because of a balanced description of core and valence electrons [136,137]. The NESCau/M06 geometry optimisations and frequencies calculations, as well as the subsequent local mode analysis was carried out with the Cologne2020 program package [138]. For the NBO analysis, the NBO 6 program [139] was used, whereas the topological electron density analysis of \( \rho(r) \) was carried out with the program AIMAll [140].

3. Results and discussion
3.1. Geometries and strength of the an-ring interaction

In order to determine the position of \( M \), the Cartesian coordinate system is rotated into the standard orientation with uniquely defined the standard x-, y- and z-axes, which is defined by the Cremer-Pople mean ring plane [123,124] of the ring atoms. As sketched in Figure 3, one can then uniquely define a projected stretching direction \( \mathbf{e}_{1,2} = \mathbf{r}_{1,2}/r_{1,2} \) along the normal vector direction \( \mathbf{n} \) of the ring’s unique mean plane [123,124], and derive the force constant \( k^A(An-M) \) and related frequency \( \omega^A(An-M) \).

The local mode analysis was complemented with the Natural Bond Orbital (NBO) analysis [105,125–127] to determine the 5f orbital occupation number of the An atom in each compound, in this way identifying which of the 5f orbitals are participating in bonding. In order to examine the covalent character of the An–C bonds we applied the Cremer-Kraka Criterium for covalent bonding, which is based on the local energy density

\[ H(r) = G(r) + V(r) \]

where \( G(r) \) is kinetic energy density (positive, destabilising) and \( V(r) \) is potential energy density (negative, stabilising). Taken at the bond critical point \( r_b \) of the electron density \( \rho(r) \) between two bonded atoms, \( H(r_b) < 0 \) indicates a covalent bond, while \( H(r_b) > 0 \) indicates an electrostatic interaction.

The optimised geometries of all complexes An\((C_nH_n)_2\) (An : Th, Pa, U, Np, Pu; \( n = 5–8 \)) are of \( D_{5h} \) symmetry with a linear M-An-M arrangement, with one exception; the An\((C_6H_6)_2\) complexes adapt \( C_{2v} \) symmetry with a bent M-An-M arrangement (see Figure 2), in line with other studies [141–144]. All complexes were identified as minimum structures on the potential energy surface; i.e. no imaginary vibrational frequencies were encountered.

Possible reasons for the bent structures are i) the competition between ligand repulsion and bond stabilisation, discussed in the literature as a common feature of actinide sandwich complexes [145,146], or ii) a Jahn-Teller or pseudo-Jahn Teller distortion [147,148], which so far has been mostly observed for 3d-metal sandwich compounds [149,150]. Work is in progress to further explore possible Jahn-Teller distortions in 5f-metal complexes in more detail.

The results of our relativistic calculations are presented in Table 1. The calculated distances between the metal and the centre of the ring (An-M) for Th\((C_5H_5)_2\), U\((C_6H_6)_2\) and Np\((C_8H_8)_2\) are 1.999, 1.930, 1.911 Å, respectively, in good agreement with the corresponding experimental values of 2.004, 1.926 and 1.909 Å [151,152]. The same holds for the Fe-M distance of Fe\((C_5H_5)_2\); the calculated distance is 1.644 Å and the experimental distance is 1.648 Å [153,154]. This confirms the validity of our chosen NESCau/M06 model chemistry.
The weakest metal–ring interactions are observed for the Np-complexes with an average $k^a(An-M)$ value of 1.200 mDyn/Å, and the strongest interactions for the Th- and Pa-complexes, with average $k^a(An-M)$ values of 2.834 and 2.600 mDyn/Å, respectively. In comparison, the ferrocene complex has a local mode force constant of 2.834 and 2.600 mDyn/Å, respectively. For the sandwich complexes with 6-membered (6MR) rings, we can see some trends. The ferrocene complex has a local mode force constant of 2.834 and 2.600 mDyn/Å, respectively. In comparison, the ferrocene complex has a local mode force constant of 2.834 and 2.600 mDyn/Å, respectively.

Although there is no overall significant correlation between distance and strength, we can see some trends. For the sandwich complexes with 6-membered (6MR) and 7-membered (7MR) rings we find a Badger-type [155] relationship, i.e. for each of the series a shorter distance relates to a stronger interaction. However, for the sandwich complexes with five-membered (5MR) rings, (longest An-M distances) and 8-membered (8MR) rings, (shortest An-M distances), we find an inverse Badger relationship, i.e. within each of this series the longer An-M distance is paired with larger local force constant. In both cases, the ring is negatively charged leading to electrostatic repulsion between ring and metal centre. This is in line with previous studies where we observed increased bond weakening with decreasing bond length for chemical bonds between two electronegative atoms (e.g. NF).
caused by increasing electrostatic repulsion, which over-
rules the Badger rule [83,156].

Figure 5 correlates the local mode force constants $k^a(An-C)$ averaged over all ring An-C bonds for each ring and the local stretching force constant $k^a(An-M)$ for the metal–ring centre interaction. There is a significant correlation between both local force constants for the 5MR, 7MR and 8 MR sandwich compounds. However, Figure 5 clearly reveals that the former are smaller, i.e. the strength of the interaction between the metal and the individual carbon atoms of the ring as bond strength measure. This is also reflected by the local vibrational frequencies $\omega^a$ values collected in Table 1. In the case of the bent 6MR sandwich complexes the correlation between the two force constants falls short, confirming that only the local $k^a(An-M)$ force constants are suited as intrinsic bond strength measure, applicable to complexes with higher and lower symmetry.

3.2. NBO analysis and 5f occupation

Actinides are characterised by gradually filling the 5f-electron shell. The level of localisation and participation of the actinide 5f valence orbitals in covalent bonds across the actinide series is still an ongoing debate [157,158]. The 5f orbitals of actinides can be localised like 4f orbitals in the lanthanide series, or delocalised like d orbitals in transition metals. Also, relativistic effects can move the 5f and 6d orbital energies closer together, allowing both orbitals to participate in chemical bonding [159].

In order to describe the role of 5f electrons in An-ring bonding, the 5f orbital occupation number of the An atom of each compound investigated in this work was calculated via an NBO analysis and compared with the corresponding local stretching force constant $k^a(An-M)$. In Table 1 all 5f occupation numbers are collected. In Figure 6, the 5f occupation numbers are compared with the local stretching force constants $k^a(An-M)$. An isolated Th atom with the electron configuration of [Xe]6d²7s² has no 5f occupation. This is reflected by the close to zero values in Table 1 found for all Th sandwich complexes, independent of the type and size of the ring. However, as depicted in Figure 6 the local $k^a(An-M)$ force constants vary from 2.361 mDyн/Å (6MR) to 3.584 mDyн/Å (8MR), showing a clear variation in metal–ring bonding. This indicates that 5f molecular orbitals do not participate in the Th-ring interaction, in line with the finding that Th tends to resemble d-block transition metals [160,161]. The 5f occupation numbers of the Pa complexes are in the range of 1.18 e (5MR) –1.73 e (8MR), being smaller than the 5f occupation number in the isolated Pa atom ([Xe]5f²6d⁰7s²). However, there is no direct correlation with the local stretching force constants $k^a(Pa-M)$, i.e. the interaction is not dominated by 5f electron participation.

The same holds for the 5f occupation numbers of the U-complexes ranging from 2.43 e (7MR) –2.81 e (5MR) which are smaller than that of the isolated U atom ([Xe]5f³6d¹7s²). Again, there no direct correlation between 5f delocalisation and bond strength, the largest $k^a(U-M)$ value of 2.417 mDyн/Å is found for the 8MR while the smallest 5f occupation number of 2.43 e, i.e.
largest delocalisation is found for the 7MR. The 5f occupation numbers for the Np-complexes are close to the configuration of the isolated Np atom ([Xe]5f⁴6d¹7s²). It is interesting to note that in the case of the 5MR and 8MR the 5f occupation number is larger than 4 (4.22 e, 5MR and 4.19 e, 8MR) indicating a delocalisation of charge from the negatively charged ring into the 5f orbitals, which correlates with the higher kₐ(An-M) values of 1.261 mDyne/Å, 5MR and 1.288 mDyne/Å, 8MR. However, overall the Np complexes have the weakest metal–ring interaction compared to other An-complexes, in line with arguments that the 5f electrons in Np are more localised and less prone to metal–ring interactions [160,162]. The 5f occupation numbers of the Pu-complexes are in a range of 4.82 e (7MR)–5.79 e (8MR) which are again smaller than the 5f occupation number in the isolated Pu atom ([Xe]5f⁶6d⁰7s²). But also in this case there is no direct correlation between 5f electron occupation and the strength of the metal–ring interaction.

3.3. Electron and energy density distribution

The local mode and the NBO analyses were complemented with the topological analysis of the electron density ρ(r) focusing on the characterisation of the An-C bonds. Table 1 collects the electron density ρ(rₜₜ) at the An-C bond critical point rₜₜ averaged over all An-C ring bonds for each ring and the corresponding energy density H(rₜₜ). For 5MR and 6MR complexes in addition to bond critical points rₜₜ also cage critical points rₜ were found. The corresponding electron density ρ(rₜₜ) and energy density H(rₜ) at the cage critical point are also included in Table 1. In Figure 7(a), ρ(rₜₜ) and ρ(rₜₜ) values are compared with the local mode force constants kₐ(An-M), whereas Figure 7(b) relates energy density H(rₜₜ) and H(rₜ) with the local mode force constants kₐ(An-M).

As discussed above, the covalent character of a chemical bond is reflected by a negative value of H(rₜₜ) while a positive energy density H(rₜₜ) corresponds to a chemical bond with dominant electrostatic character [129,130,163]. The energy density H(rₜₜ) of the Th-complexes is between −0.0410 and −0.0495 Hartree/Å³ indicating the covalent character of the Th-C ring bonds, which correlates with larger local mode force constants kₐ(An-M), see Table 1 and Figure 7(b). Pa(C₆H₆)₂, Pa(C₇H₇)₂ and Pa(C₈H₈)₂ complexes have H(rₜₜ) values from −0.0021 to 0.0045 Hartree/Å³, showing a mixture of the electrostatic and the covalent bonding between metal and ring atoms. The Pa(C₆H₆)₂ complex which has a bent geometry contrary to the linear M–Pa–M arrangement of the other Pa sandwich complexes, has an H(rₜₜ) value of −0.0304 Hartree/Å³, revealing larger covalent character of the Pa–C bonds. This can be explained by the larger overlap between the π orbitals of benzene with the 6d and 5f orbitals of the metal in the bent geometry, leading to stronger electron donation from the ring and to more covalent bonding character. The same effect is true for the bent U(C₆H₆)₂, Np(C₆H₆)₂ and Pu(C₆H₆)₂ complexes. The only exception is the bent Th(C₆H₆)₂ complex, showing less covalent character than the other Th complexes. In this case, the 5f orbitals are empty, and the bent structure does not necessarily lead to a stronger electron donation from the ring to the metal atom. Overall, as depicted in Figure 7(a,b), there is no direct correlation between An-C bond critical point properties and the intrinsic strength of the An-ring interaction represented via the local mode force constants kₐ(An-M).

4. Conclusion

This work provides a detailed analysis of An(CₙHₙ)₂ (An: Th, Pa, U, Np, Pu; n = 5–8) sandwich complexes combining the NESCaU approach and the local vibrational mode analysis. We could shed new light on
the An-carbon bonding and the An-ring interactions in these f-block sandwich complexes including the following highlights:

- Our study shows that NESCau is an easily applicable method providing a solid fundament for the calculation of compounds with superheavy elements. Building upon the locality of relativistic effects, NESCau makes NESC calculations dozens to hundreds of times faster without losing accuracy [56].
- An(C₇H₁₅)₂ complexes with smaller ring are characterised by longer An-ring distances than complexes with larger rings. However, the strength of the An-ring interaction does not correlate with ring size and/or metal–ring distance; it is determined by the character of the metal centre. The weakest An-ring interactions were observed for Np(C₇H₁₅)₂ while the strongest for Th(C₇H₁₅)₂ complexes.
- Calculated 5f occupation numbers are in line with the expected An oxidation states. They increase across the actinide series denoting a successive increase of the electron localisation on the An centre with increasing atomic number. As reflected by the corresponding local mode force constants, delocalisation of the 5f-orbitals leads to a weakening of the An-ring interaction. Th complexes with a 5f occupation number close to zero have the strongest An-ring interactions. This clearly shows that in these compounds 5f orbitals do not play a role for the metal–ring interactions.
- According to the Cremer-Kraka criterion strongest covalent An-C bonds were observed in Th-complexes, followed by U and Np complexes, while Pu and in particular Pa complexes showed a mixture of covalent bonding and electrostatic interactions.

By rationalising the intrinsic strength of An-C bonding and An-ring interactions and their interplay in a series of 20 An sandwich complexes An(C₇H₁₅)₂, this work provides valuable information for the fine-tuning and future synthesis of An sandwich compounds. We also hope to encourage by our study the use of NESCau for the investigation of large systems with heavy metal and superheavy metal atoms.

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