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The density-based many-body expansion for poly-peptides and proteins†

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Fragmentation schemes enable the efficient quantum-chemical treatment of large biomolecular systems, and provide an ideal starting point for the development of accurate machine-learning potentials for proteins. Here, we present a fragment-based method that only uses calculations for single-amino acids and their dimers, and is able to reduce the fragmentation error in total energies to ca. 1 kJ mol⁻¹ per amino acid for polypeptides and proteins across different structural motifs. This is achieved by combining a two-body extension of the molecular fractionation with conjugate caps (MFCC) scheme with the density-based many-body expansion (db-MBE), thus extending the applicability of the db-MBE from molecular clusters to polypeptides and proteins.

1 Introduction

Molecular dynamics simulations make it possible to study the structural dynamics of biomolecules in solution, and provide valuable insights into the emergence and mechanisms of biological function. 1,2 Conventionally, such simulations rely on classical force fields to model the potential energy surface of biomolecules such as proteins.³⁻⁷ In the past decade, considerable efforts were undertaken to extend the time scales of classical molecular dynamics simulations, which made it possible to directly simulate protein folding processes.^{8–10}

However, the accuracy of classical force fields is inherently limited, 11,12 and the steep increase in computational effort of more accurate quantum-chemical methods makes them rarely applicable to large biomolecules. 13,14 For treating excited state phenomena in biological systems^{15,16} and to study excited-state dynamics, ¹⁷⁻²⁰ quantum-chemical methods are mandatory. Fragmentation methods²¹⁻³⁰ provide an avenue to decreasing the computational effort of quantum-chemical calculations and to lower its scaling with system size. This is achieved by replacing a calculation for the full system (e.g., a protein) by many small calculations for its fragments (e.g., single amino acids).

Recently, such quantum-chemical fragmentation schemes have been used as reference for the parametrization of machine-learning potentials for polypeptides and proteins. These methods hold the promise of enabling molecular dynamics simulations on a

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highly accurate potential energy surface, possibly matching the accuracy of high-level quantum chemistry, and even allow for extensions to excited state dynamics. 35-37 However, any machine learning potential can only be as good as the quantum-chemical reference data on which it has been trained. This includes not only the underlying quantum-chemical methods, but also the accuracy of the fragmentation scheme.

Therefore, the development of quantum-chemical fragmentation schemes remains an important research avenue (for recent efforts, see, e.g., ref. 38-49). Such fragmentation schemes should be (a) versatile in both their applicability to complex chemical systems, including proteins, and in their compatibility with quantum-chemical methods for the fragment calculations, (b) only introduce a small error due to the fragmentation while maintaining a reasonable computational cost, and (c) ideally only use a small number of chemically meaningful fragments. In the past years, our research group has pursued two lines of research towards this goal (see Fig. 1).

First, for systems composed of distinct molecular building blocks such als molecular clusters, we have extended the conventional energy-based many-body expansion (eb-MBE)50-52 by a density-based correction, that is derived by performing a manybody expansion of the electron density, and inserting the resulting total density into the total energy functional of density-functional theory (DFT).⁵³ This density-based many-body expansion (db-MBE) can be considered as an ONIOM-style54,55 multilevel method, in which the high-level is provided by the eb-MBE and the low-level is provided by subsystem-DFT. 56,57 Previously, we have demonstrated that for water and ion-water clusters, the db-MBE provides accurate and efficient total and relative energies already at the level of a two-body expansion. 58-60

Second, for proteins we have extended the (energy-based) molecular fractionation with conjugate caps (eb-MFCC) method

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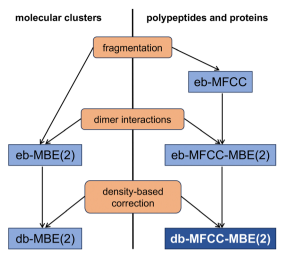


Fig. 1 Overview of the relationship between the different fragmentation schemes considered in this work, up to second order. Schemes for molecular clusters are shown on the left, while the corresponding schemes for polypeptides and proteins are given on the right. The boxes in the middle specify the additional contributions included in each of these schemes. The db-MFCC-MBE(2) scheme is the topic of this paper.

to consistently include two-body (dimer) contributions [eb-MFCC-MBE(2)].61 While our eb-MFCC-MBE(2) scheme can be considered as a special case of more general schemes for energy-based manybody expansions with overlapping fragments, 41,62,63 it provided a simple and consistent approach for the quantum-chemical calculation of the energy of proteins and of protein-ligand interaction energies.⁶⁴ The eb-MFCC and eb-MFCC-MBE(2) schemes are particularly attractive as starting point for the parametrization of machine-learning potentials because they use single amino acids as fragments.31,32

However, our tests showed^{61,64} that while the inclusion of two-body contributions in the MFCC-MBE(2) scheme dramatically reduced the error in protein energies and protein-ligand interaction energies compared to the MFCC scheme, the remaining errors can still be substantial.

In the present work, we thus combine the eb-MFCC-MBE(2) approach with the db-MBE, thereby extending the scope of the db-MBE from molecular clusters to proteins. We apply the resulting db-MFCC-MBE(2) to various test cases and assess its accuracy for total energies of polypeptides and proteins as well as for relative energies of polypeptide conformers.

Computational methodology

eb-MBE and db-MBE

First, we briefly review the eb-MBE, 50-52,65-68 considering a cluster consisting of N molecular fragments. Using an n-body expansion, its total energy is approximated as

$$E_{\text{tot}} \approx E_{\text{eb}}^{(n)} = E_{\text{eb}}^{(1)} + \sum_{m=2}^{n} \Delta E_{\text{eb}}^{(m)},$$
 (1)

where the first-order term

$$E_{\rm eb}^{(1)} = \Delta E_{\rm eb}^{(1)} = \sum_{i_1}^{N} E_{i_1}.$$
 (2)

is the sum of the energies E_{i} , of the monomers. The *n*-th order term is given by

$$\Delta E_{eb}^{(n)} = \sum_{i_1=1}^{N} \sum_{i_2=i_1+1}^{N} \dots \sum_{i_n=i_{n-1}+1}^{N} E_{i_1 i_2 \dots i_n} - \sum_{m=1}^{n-1} \frac{N^{n-m}}{(n-m)!} \cdot \Delta E_{eb}^{(m)},$$
(3)

where E_{i,i,\dots,i_n} is the energy of the *n*-mer consisting of molecular fragments $\{i_1,i_2,...,i_n\}$. For instance, the second-order (twobody) term is given by

$$\Delta E_{eb}^{(2)} = \sum_{i_1=1}^{N} \sum_{i_2=i_1+1}^{N} E_{i_1 i_2} - N \sum_{i_1}^{N} E_{i_1}$$

$$= \sum_{i_1=1}^{N} \sum_{i_2=i_1+1}^{N} (E_{i_1 i_2} - E_{i_1} - E_{i_2}). \tag{4}$$

In the same fashion as the total energy, the electron density can be approximated using a many-body expansion as

$$\rho_{\text{tot}}^{(n)} = \rho_{\text{tot}}^{(1)} + \sum_{m=2}^{n} \Delta \rho_{\text{tot}}^{(m)}, \tag{5}$$

with the first-order approximation

$$\rho_{\text{tot}}^{(1)} = \Delta \rho_{\text{tot}}^{(1)} = \sum_{i_1}^{N} \rho_{i_1}, \tag{6}$$

and the n-th-order density correction,

$$\Delta \rho_{\text{tot}}^{(n)} = \sum_{i_1=1}^{N} \sum_{i_2=i_1+1}^{N} \dots \sum_{i_n=i_{n-1}+1}^{N} \rho_{i_1 i_2 \dots i_n} - \sum_{m=1}^{n-1} \frac{N^{n-m}}{(n-m)!} \cdot \Delta \rho_{\text{tot}}^{(m)}.$$
 (7)

Here, ρ_{i_1} is the electron density of the i_1 -th monomer, and $\rho_{i,i,\ldots i_n}$ is the electron density of the *n*-mer consisting of molecular fragments $\{i_1,i_2,\ldots,i_n\}$. In analogy to the many-body expansion of the total energy [see eqn (4)], the second-order (two-body) correction to the electron density is given by

$$\Delta \rho_{\text{tot}}^{(2)} = \sum_{i_1=1}^{N} \sum_{i_2=i_1+1}^{N} \rho_{i_1 i_2} - N \sum_{i_1}^{N} \rho_{i_1}$$

$$= \sum_{i_1=1}^{N} \sum_{i_2=i_1+1}^{N} \left(\rho_{i_1 i_2} - \rho_{i_1} - \rho_{i_2} \right). \tag{8}$$

Other semi-local ingredients (such as the density gradient, density Hessian, or the kinetic energy density) can be approximated in the same fashion as the electron density.

Using such a many-body expansion of the electron density, the *n*th-order db-MBE is then defined as, 58

$$E_{\rm db}^{(n)} = E[\rho_{\rm tot}^{(n)}] = E_{\rm eb}^{(n)} + \Delta E_{\rm db-eb}^{(n)}, \tag{9}$$

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where $E[\rho]$ is the DFT total energy functional and the densitybased correction is at *n*-th order given by

$$\Delta E_{\text{db-eb}}^{(n)} = E[\rho_{\text{tot}}^{(n)}] - E_{\text{eb}}^{(n)}.$$
 (10)

This density-based correction can be evaluated using only the electron densities from the fragment calculations (i.e., without explicit reference to the energy contributions entering $E_{\rm eh}^{(n)}$, and is given by⁵⁸

$$\begin{split} \Delta E_{\text{db-eb}}^{(n)}[\rho_{i_{1}},\rho_{i_{1}i_{2}},\dots] &= \left(V_{\text{nuc}}[\rho_{\text{tot}}^{(n)}] - V_{\text{nuc}}^{(n)}\right) + \left(J[\rho_{\text{tot}}^{(n)}] - J^{(n)}\right) \\ &+ \left(E_{\text{NN}} - E_{\text{NN}}^{(n)}\right) + T_{\text{s}}^{\text{nadd},(n)}[\rho_{i_{1}},\rho_{i_{1}i_{2}},\dots] \\ &+ E_{\text{xc}}^{\text{naddd},(n)}[\rho_{i_{1}},\rho_{i_{1}i_{1}},\dots], \end{split} \tag{11}$$

with the *n*-body nonadditive kinetic and exchange-correlation energy functionals defined as

$$T_{\rm s}^{{\rm nadd},(n)}[\rho_{i,},\rho_{i,i,},\dots] = T_{\rm s}[\rho_{\rm tot}^{(n)}] - T_{\rm s}^{(n)}$$
 (12)

$$E_{\text{xc}}^{\text{nadd,(n)}}[\rho_{i_1}, \rho_{i_1 i_2}, \dots] = E_{\text{xc}}[\rho_{\text{tot}}^{(n)}] - E_{\text{xc}}^{(n)},$$
 (13)

which are evaluated using approximate kinetic energy and exchange-correlation density functionals, akin to subsystem DFT. 56,57 In these expression, $V_{\text{nuc}}^{(n)}$, $J_{\text{nuc}}^{(n)}$, $E_{\text{NN}}^{(n)}$, $T_{\text{s}}^{(n)}$, and $E_{\text{xc}}^{(n)}$ are the n-body expansions of the individual contributions to the DFT total energy functional, which are defined in analogy to eqn (1). Note that a correction due to the nuclear repulsion energy only appears at first order, because $E_{NN}^{(n)} = E_{NN}$ for $n \ge 2$. Explicit expressions for $\Delta E_{\text{db-eb}}^{(n)}$ at the one-body and two-body levels are presented in Section S1.1 of the ESI.†

2.2 eb-MFCC and eb-MFCC-MBE(2)

The MFCC scheme⁶⁹⁻⁷¹ has been devised to provide a simple approach for approximating the total energies and electron densities of proteins and to calculate approximate protein-ligand interaction energies. 72-76 In its simplest form, the protein is separated into single amino acids by placing cuts across the peptide bonds. The unsaturated bonds are then capped by acetyl (ACE) and N-methylamide groups (NME) (see Fig. 2). Disulfide bridges are similarly cut and capped by methyl sulfide groups.

The total energy of a system consisting of N amino acids is then approximated as the sum of all capped fragment energies $E_{\rm eb}^{\rm f}$ from which the sum of all cap molecule energies $E_{\rm eb}^{\rm c}$ is subtracted, leading to the eb-MFCC energy expression

$$E_{\text{tot}}^{\text{MFCC}} = E_{\text{eb}}^{(1)} = E_{\text{eb}}^{\text{f}} - E_{\text{eb}}^{\text{c}} = \sum_{i_1=1}^{N} E_{i_1}^{\text{f}} - \sum_{k_1=1}^{N-1} E_{[k_1,k_1+1]}^{\text{c}}, \quad (14)$$

where $E_{i_1}^{\rm f}$ is the total energy of the i_1 -th capped fragment and $E_{[k_1,k_1+1]}^{c}$ is the total energy of the ACE-NME molecule formed from the caps of fragments k_1 and $k_1 + 1$. For consistency with the above expressions for the MBE and notational simplicity, we drop the superscript MFCC and refer to the above first-order approximation for the total energy as $E_{\rm eb}^{(1)}$.

The most important shortcoming of the MFCC scheme is its neglect of intramolecular interactions. Several extensions of the MFCC scheme have been proposed to alleviate this issue⁷⁷⁻⁸¹ (see also ref. 61 and references therein). One approach for systematically improving the MFCC scheme is its combination with the many-body expansion. To include pairwise interactions (i.e., two-body contributions), fragment-fragment (ff) interaction energies are calculated akin to the second-order eb-MBE. To take account of the accruing interactions between fragments and caps as well as interactions between caps, one additionally needs to account for fragment-cap (fc) and capcap (cc) interactions. The resulting eb-MFCC-MBE(2) energy expression reads,61

$$E_{\text{tot}}^{\text{MFCC}} = E_{\text{eb}}^{(2)} = E_{\text{eb}}^{(1)} + \Delta E_{\text{eb}}^{(2)}$$
 (15)

with the two-body correction

$$\Delta E_{\rm eb}^{(2)} = \Delta E_{\rm tot}^{\rm MFCC\text{-}MBE(2)} = \Delta E_{\rm eb}^{\rm ff} - \Delta E_{\rm eb}^{\rm fc} + \Delta E_{\rm eb}^{\rm cc}$$
 (16)

Here, the fragment-fragment contributions are given by

$$\Delta E_{\rm eb}^{\rm ff} = \sum_{i_1=1}^{N} \sum_{i_2=i_1+1}^{N} \Delta E_{i_1 i_2}^{\rm ff}, \tag{17}$$

fragment-fragment interaction energies are where the calculated as

$$\Delta E_{i_1 i_2}^{\rm ff} = \begin{cases} E_{i_1, i_1 + 1}^{\rm ff} - \left[E_{i_1}^{\rm f} + E_{i_1 + 1}^{\rm f} - E_{[i_1, i_1 + 1]}^{\rm c} \right] & \text{for } i_2 = i_1 + 1 \\ E_{i_1, i_1 + 1, i_1 + 2}^{\rm fff} - \left[E_{i_1, i_1 + 1}^{\rm ff} + E_{i_1 + 1, i_1 + 2}^{\rm ff} - E_{i_1 + 1}^{\rm f} \right] & \text{for } i_2 = i_1 + 2 \\ E_{i_1 i_2}^{\rm ff} - \left[E_{i_1}^{\rm f} + E_{i_2}^{\rm f} \right] & \text{for } i_2 > i_1 + 2 \end{cases}$$

$$(18)$$

that is, applying different formulas for neighboring $(i_2 = i_1 + 1)$ and distant $(i_2 > i_1 + 2)$ fragment pairs as well as an indirect formulation for next-nearest neighbor pairs ($i_2 = i_1 + 2$), in which the cap molecule would overlap with the capped fragments. The fragment-cap contributions are given by

$$\Delta E_{\text{eb}}^{\text{fc}} = \sum_{i_1=1}^{N} \sum_{\substack{k_1=1\\k_1 \neq i_1 - 2, \dots, i_1 + 1}}^{N-1} \Delta E_{i_1, [k_1, k_1 + 1]}^{\text{fc}}, \tag{19}$$

with the fragment–cap interaction energies $\Delta E^{\rm fc}_{i_1,[k_1,k_1+1]} = E^{\rm fc}_{i_1,[k_1,k_1+1]} - E^{\rm f}_{i_1} - E^{\rm c}_{[k_1,k_1+1]}$ (where $E^{\rm fc}_{i_1,[k_1,k_1+1]}$ is the total energy

$$\mathsf{E} \left(\mathsf{H}_2 \mathsf{N} + \mathsf{D} \right) = \mathsf{E} \left(\mathsf{H}_2 \mathsf{N} + \mathsf{D} \right) + \mathsf{E} \left(\mathsf{D} \right) - \mathsf{E} \left(\mathsf{D} \right) - \mathsf{E} \left(\mathsf{D} \right) + \mathsf{D} \right)$$

Fig. 2 Illustration of the MFCC partitioning scheme for an alanine dipeptide. The peptide bond is cut and the resulting fragments are capped with Nmethylamide groups (blue) and acetyl groups (red). A new cap molecule N-methylacetamide is then formed by the combination of both caps.

of the dimer formed from fragment i_1 and cap $[k_1,k_1+1]$). Note that caps neighboring the capped fragments are excluded from the summation by excluding caps with indices $k_1=i_1-2,\ldots,i_1+1$ due to overlapping caps. Finally, the cap–cap contributions are given by

$$\Delta E_{\rm cb}^{\rm cc} = \sum_{k_1=1}^{N-1} \sum_{k_2=k_1+2}^{N-1} \Delta E_{[k_1,k_1+1][k_2,k_2+1]}^{\rm cc}, \tag{20}$$

with the cap-cap interaction energy $\Delta E^{\rm cc}_{[k_1,k_1+1][k_2,k_2+1]}=E^{\rm cc}_{[k_1,k_1+1][k_2,k_2+1]}-E^{\rm c}_{[k_1,k_1+1]}-E^{\rm c}_{[k_2,k_2+1]}$. Further details can be found in ref. 61, and an extension of the eb-MFCC-MBE(2) scheme to protein-ligand interaction energies has been presented in ref. 64.

2.3 db-MFCC and db-MFCC-MBE(2)

The MFCC and MFCC-MBE(2) schemes can straightforwardly be extended to the electron density. At first order, the electron density can be approximated as,

$$\rho_{\text{tot}}^{(1)} = \rho_{\text{tot}}^{\text{MFCC}} = \rho_{\text{tot}}^{\text{f}} - \rho_{\text{tot}}^{\text{c}} = \sum_{i_1=1}^{N} \rho_{i_1}^{\text{f}} - \sum_{k_1=1}^{N-1} \rho_{[k_1,k_1+1]}^{\text{c}}$$
(21)

and the MFCC-MBE(2) second-order correction to the electron density can be calculated as

$$\Delta \rho_{\text{tot}}^{(2)} = \Delta \rho_{\text{tot}}^{\text{ff}} - \Delta \rho_{\text{tot}}^{\text{fc}} + \Delta \rho_{\text{tot}}^{\text{cc}} \tag{22}$$

with

$$\Delta \rho_{\text{tot}}^{\text{ff}} = \sum_{i_1=1}^{N} \sum_{i_2=i_1+1}^{N} \Delta \rho_{i_1 i_2}^{\text{ff}}$$
 (23)

$$\Delta \rho_{\text{tot}}^{\text{fc}} = \sum_{i_1=1}^{N} \sum_{\substack{k_1=1\\k_1 \neq i_1-2,\dots,i_1+1}}^{N} \Delta \rho_{i_1,[k_1,k_1+1]}^{\text{fc}}$$
(24)

$$\Delta \rho_{\text{tot}}^{\text{cc}} = \sum_{k_1=1}^{N-1} \sum_{k_2=k_1+2}^{N-1} \Delta \rho_{[k_1,k_1+1],[k_2,k_2+1]}^{\text{cc}}.$$
 (25)

In these expressions, all terms are defined in complete analogy to the respective energy terms.

By inserting these expansions of the electron density into eqn (9) and (10), we arrive at the energy expressions for the corresponding density-based variants. The db-MFCC energy is given by,

$$E_{\rm db}^{(1)} = E_{\rm tot}^{\rm db-MFCC} = E_{\rm eb}^{(1)} + \Delta E_{\rm db-eb}^{(1)},$$
 (26)

and the db-MFCC-MBE(2) energy is

$$E_{\rm db}^{(2)} = E_{\rm tot}^{\rm db-MFCC-MBE(2)} = E_{\rm eb}^{(2)} + \Delta E_{\rm db-eb}^{(2)}.$$
 (27)

The density-based correction can be evaluated using eqn (11), where for the many-body expansions of the individual energy terms, the more general definition including the contributions of the cap is used. We note that the first-order density-based correction is equivalent to the interaction energy appearing in the generalization of subsystem DFT to the MFCC

partitioning. 81 Further details are given in Section S1.2 of the ESI. †

2.4 Implementation and computational details

The db-MFCC-MBE(2) method as described above has been implemented in the PyADF scripting framework⁸² and its PyEmbed module,⁸³ based on the existing implementations of the eb-MFCC-MBE(2) scheme⁶¹ and of the db-MBE.^{53,58} The implementation follows what has been described in our previous publications. The source code of PyADF 1.5, which is suitable for all calculations presented in this work, is available as open source software at ref. 84.

The structures of all considered proteins have been obtained from the protein data bank. Hydrogen atoms were added using OpenBabel, ^{85,86} considering a neutral protonation state for each amino acids. In the process of partitioning the proteins, ACE-NME caps are added using the original positions for all cap atoms that are already present in the protein, while the remaining hydrogen atoms of the methyl groups are added using a fixed C–H bond distance of 1.07 Å. Disulfide bridges between amino acids are capped with methylsulfide groups. The resulting dimethyldisulfide cap molecules are treated analogously to the other cap molecules in the many-body corrections.

All quantum-chemical calculations have been performed using density-functional theory (DFT) with the Amsterdam Density Functional (ADF) program⁸⁷ in the Amsterdam Modeling Suite (AMS)⁸⁸ with the BP86 exchange-correlation functional^{89,90} and a DZP basis set.⁹¹ In all calculations, we used a Becke integration grid of "normal" accuracy.⁹² All total energies have been obtained with ADF's total energy implementation.⁹³ These technical settings have previously been tested in the context of the db-MBE.^{53,58}

In the eb-MFCC-MBE(2) and db-MFCC-MBE(2) calculations, a distance cut-off of 4 Å has been used, *i.e.*, calculations for dimers that are further apart than this cutoff are skipped. The choice of the cut-off is based on our previous tests in ref. 61. The evaluation of the different terms in the density-based energy correction for db-MFCC and db-MFCC-MBE(2) has been implemented as described in ref. 58. All these terms have been evaluated using the supermolecular numerical integration grid. In the evaluation of the Coulomb contributions, the corrections necessary for ADF's fitted density are included consistently. Again, these settings have previously shown to be adequate for db-MBE calculation of molecular clusters. The revaluating the nonadditive exchange-correlation and kinetic energy functionals, the XCFun library stream was used, and the BP86 and PW91k⁹⁷ functionals were applied.

3 Results and discussion

3.1 Alanine polypeptides

As our initial test case, we consider four alanine polypeptides as idealized models for different secondary structure elements. The test set comprises (Ala)₁₀ in a 3₁₀-helix conformation, as an α -helix, and as a β -strand, as well as (Ala)₁₁ featuring a turn between two antiparallel strands, as model of a β -sheet (see insets in Fig. 3). Each of these models was constructed by

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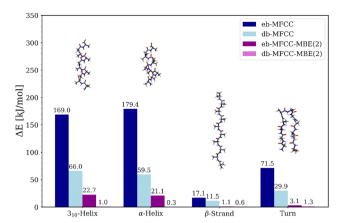


Fig. 3 Comparison of the absolute errors in the total energy (DFT/BP86/ DZP) with the energy-based and the density-based MFCC and MFCC-MBE(2) schemes for (Ala) $_{10}$ in a 3_{10} -helix conformation, as an α -helix, and as a β -strand as well as for an (Ala)₁₁ turn. As reference, single-point calculations for the full polypeptides have been performed.

assuming idealized backbone angles for the respective secondary structure element. These model polypeptides were previously considered in ref. 61.

In Fig. 3, the absolute errors in the total energy compared to a supermolecular DFT calculation are plotted for both the eb-MFCC and eb-MFCC-MBE(2) schemes and for the corresponding density-based extensions of these schemes. In all cases, the conventional eb-MFCC scheme leads to rather large errors. For the two helices, the errors of 169 and 179 kJ mol⁻¹ demonstrate the need of accounting for intramolecular interactions. For the turn, the error is smaller because there are fewer intramolecular hydrogen bonds, but with 72 kJ mol⁻¹, it is still substantial. Even for the β-strand, in which no intramolecular hydrogen bonds are present, the error still amounts to 17 kJ mol^{-1} .

Including two-body contributions in the eb-MFCC-MBE(2) reduces these error significantly (see also ref. 61). For the β -strand and the turn, this is sufficient to reduce the errors to 1.1 and 3.1 kJ mol⁻¹, respectively, whereas for the 3_{10} -helix and the α -helix, errors of 22.7 and 21.1 kJ mol⁻¹, respectively, remain. In the latter case, each peptide group in the center of the helix is involved in two hydrogen bonds, such that these intramolecular interactions are not fully captured by a two-body approximation.

For both MFCC and MFCC-MBE(2), adding the densitybased correction lowers the error significantly. For the db-MFCC scheme, the errors for the 3_{10} -helix and the α -helix are reduced to 66 and 60 kJ mol⁻¹, respectively, while for the β-strand and the turn, they are reduced to 12 and 30 kJ mol⁻¹, respectively. When including two-body contributions in the db-MFCC-MBE(2) scheme, the calculations become virtually exact, with errors between 0.3 and 1.3 kJ mol⁻¹ in the total energies. This demonstrates that the density-based correction is able to account for higher-order contributions, that would only appear at third or higher orders in the energy-based expansion.

3.2 Proteins

As a next step, we applied the methods previously tested to a broad range of proteins, which we assembled by considering test cases studied previously with quantum-chemical fragmentation methods^{73,98,99} as well as a search of the protein data bank for suitable proteins with a size that is still amenable to a supermolecular DFT calculation.

Our test set includes human insulin (PDB-code 3I40, 100 51 amino acids, 784 atoms), SEM5 SH3 domain (PDB-code 3SEM, 101 58 amino acids, 945 atoms), the C-terminal domain of the ribosomal protein L7/L12 (PDB code 1CTF. 102 68 amino acids, 1005 atoms), the C-terminal domain of RecA protein (PDB-code 1AA3, 103 63 amino acids, 1017 atoms), ubiquitin (PDB-code 1UBQ, ¹⁰⁴ 76 amino acids, 1231 atoms), the FADD (Mort1) death-effector domain (PDB-code 1A1W, 105 83 amino acids, 1363 atoms), the immunophilin immunosuppressant complex FKBP-FK506 (PDB code 1FKF, 106 107 amino acids, 1662 atoms), the MTCP-1 protein involved in T-cell malignancies (PDB-code 1A1X, 107 106 amino acids, 1742 atoms), and the catalytic core domain of FIV integrase (PDB-code 4PA1, 108 151 amino acids, 2378 atoms). Of these test cases, 3I40, 3SEM, 1CTF, 1UBQ, 1FKF, and 4PA1 have been used previously by us in ref. 61 to assess the accuracy of the energy-based MFCC and MFCC-MBE(2) schemes, whereas 1AA3, 1A1W, and 1AX1 have been added here.

For this test set of proteins, Fig. 4 plots the absolute errors per amino acid in the total energy compared to supermolecular calculations for the energy-based and density-based MFCC and MFCC-MBE(2) schemes. For better comparison, the errors have been normalized to the number of amino acids in the proteins. The corresponding plot of the unnormalized absolute errors is shown in Fig. S1 in the ESI.†

In agreement with our results for smaller polypeptides, the eb-MFCC scheme leads to unacceptably large errors. For 1AA3 and 1CTF, the error amounts to 42 and 18 kJ mol⁻¹ per amino acid, while for 3I40, 1UBQ, 1A1W, 1FKF, 1A1X, and 4PA1 we find errors between 5 and 14 kJ mol⁻¹ per amino acid. For 3SEM, the error is close to zero, which must be due to fortunate error cancelation in this specific case. As already observed previously, 61 the inclusion of two-body contributions in the

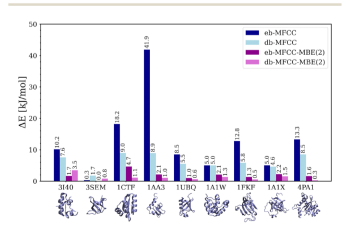


Fig. 4 Comparison of the absolute errors per amino acid in the total energy (DFT/BP86/DZP) with the energy-based and the density-based MFCC and MFCC-MBE(2) schemes for our test set of proteins. As reference, single-point calculations for the full proteins have been performed.

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eb-MFCC-MBE(2) scheme is able to reduce the error significantly, and we find errors between 1.0 and 4.7 kJ mol⁻¹ per amino acid, with the exception of 3SEM, for which the error remains close to zero.

The inclusion of the density-based correction in the db-MFCC scheme is able to reduce the errors substantially compared to the eb-MFCC scheme, resulting in errors per amino acid between 5 and 9 kJ mol⁻¹, again with the exception of 3SEM. While this is a substantial improvement—in particular when considering that no additional fragment calculations are required for evaluation the density-based correction on top of the eb-MFCC—the remaining errors show that the db-MFCC can only partially capture the twobody and higher-order interactions.

However, when including two-body contributions in the db-MFCC-MBE(2) scheme, the errors per amino acid are reduced to below 1.5 kJ mol⁻¹. In most cases, this amounts to a reduction of the error by roughly a factor of two, but in cases such as 1CTF, where the error of the eb-MFCC-MBE(2) was particularly large, even larger reductions are observed. In all cases, the absolute error per amino acid is below the threshold of chemical accuracy $(4 \text{ kJ mol}^{-1}).$

Notably, for 3SEM, for which the eb-MFCC-MBE(2) scheme already resulted in an error of close to zero, the error increases when including the density-based correction. This is consistent with the assumption that the excellent performance of the eb-MFCC and eb-MFCC-MBE(2) schemes in this particular case is due to fortunate error cancellation. Finally, we note that for the db-MFCC-MBE(2) scheme, the largest error per amino acid (3.5 kJ mol⁻¹) is found for 3I40, where the error also increases compared to the eb-MFCC-MBE(2) scheme. Again, this seems to be due to fortunate error cancellation in the energy-based expansions. Likely, the inclusion of additional two-body terms (that are neglected due to distance-based screening) and/or of three-body contributions will rectify this.

Overall, we find that the inclusion of the density-based correction on top of the eb-MFCC-MBE(2) scheme, which generally reduces the error in the total energies of the considered proteins, and for most test cases leads to an agreement with full supermolecular calculations within ca. 1 kJ mol⁻¹ per amino acid.

3.3 Relative energies for polypeptides

While the above test cases only consider a single point on the potential energy surface (i.e., a fixed protein structure), molecular dynamics simulations require a balanced description of the low-energy regions of the potential energy surface. This is particularly relevant if quantum-chemical fragmentation methods are to be used for the parametrization of machine learning potentials with the aim to replace classical biomolecular force

To assess the accuracy of the considered energy-based and density-based fragmentation schemes across different regions of the potential energy surface, we consider relative energies of three different sets of conformers, specifically (Ala)₁₀ as α -helix, (Ala)₁₁ as a turn (modeling a β -sheet structure), and (Ala)₁₀ as a β -strand. For each set, 11 (for the α -helix and the turn) or 9 (for the β -strand) snapshots extracted from molecular dynamics simulations with a classical force field (see ref. 109 and 110 for details) have been used.

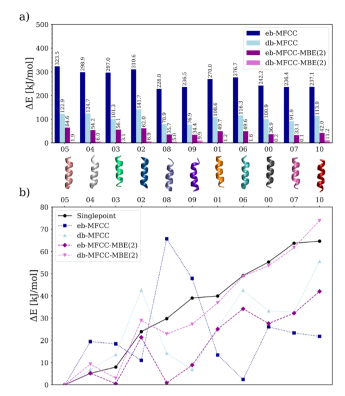


Fig. 5 Comparison of the energy-based and the density-based MFCC and MFCC-MBE(2) schemes for a test set of eleven conformers of α -helical (Ala)₁₀. (a) Absolute errors in the total energy (DFT/BP86/DZP) compared to a single-point calculations for the full polypeptides. (b) Relative energies (DFT/ BP86/DZP) (i.e., energy difference to the lowest-energy structure) of the considered conformers, ordered by increasing energy. The supermolecular reference calculations are plotted as solid black line ("Singlepoint").

Fig. 5(a) compares the absolute errors in the total energies for eleven conformers of α -helical (Ala)₁₀. The eb-MBFCC and eb-MFCC-MBE(2) schemes lead to large absolute errors that vary widely across the different conformers. With eb-MFCC-MBE(2) we find absolute errors between 35 and 65 kJ mol^{-1} . However, the inclusion of the density-based correction in the db-MFCC-MBE(2) scheme is able to reduce the error consistently, with the largest errors (9.9 and 11.2 kJ mol⁻¹) found for conformers 09 and 10.

The relative energies of the conformers are visualized in Fig. 5(b). Here, it is obvious that neither the eb-MFCC nor the db-MFCC scheme can reproduce the energy pattern correctly. The eb-MFCC-MBE(2) scheme shows a better trend, but still gets many relative energies wrong. The db-MFCC-MBE(2) scheme is able to rectify these shortcomings and closely follows the trend of the single-point reference calculations. Nevertheless, there are still some outliers, in particular for conformers 09 and 10 that also showed the largest absolute errors. Here, it is interesting to note that while for the other schemes there is significant error cancellation when considering the relative energies (i.e., all errors in the total energies have the same sign), this is not the case for the db-MFCC-MBE(2) results, where positive and negative errors in the total energies amplify when comparing relative energies.

For the $(Ala)_{11}$ turn, the absolute errors in the total energies for eleven conformers are compared in Fig. 6(a). Compared to the α -helical conformers, these absolute errors are significantly smaller, since there are fewer intramolecular interactions. Once two-body contributions are included the absolute errors are below 15 kJ mol⁻¹ for all conformers, both with eb-MFCC-MBE(2) and with db-MFCC-MBE(2). While for some conformers, the density-based scheme is able to reduce the error to close to zero, for other conformers it yields errors comparable to or even slightly worse than the energy-based scheme.

In the plot of the corresponding relative energies of the conformers in Fig. 6(b), we recognize that the eb-MFCC and db-MFCC cannot reproduce the energy ordering of the conformers correctly, even though they partly follow the correct trend. The relative energies from the eb-MFCC-MBE(2) and db-MFCC-MBE(2) schemes both closely follow the supermolecular reference, with a visually better agreement for the energy-based scheme. The db-MFCC-MBE(2) are shifted to higher relative energies, which is to a large part caused by the calculation for the lowest-energy conformer, and shows outliers for conformers 00, 01, and 02.

Finally, Fig. 7 compares the absolute errors in the total energies as well as the relative energies for nine conformers of a (Ala)₁₀ β -strand. Here, the differences in the absolute errors

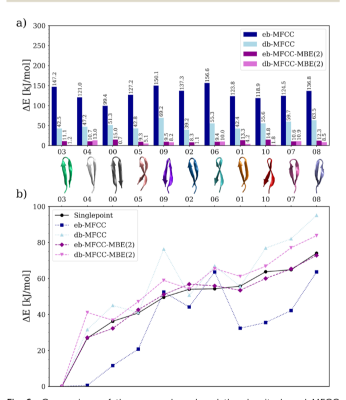


Fig. 6 Comparison of the energy-based and the density-based MFCC and MFCC-MBE(2) schemes for a test set of eleven conformers of the (Ala)₁₁ turn structure. (a) Absolute errors in the total energy (DFT/BP86/ DZP) compared to a single-point calculations for the full polypeptides. (b) Relative energies (DFT/BP86/DZP) (i.e., energy difference to the lowestenergy structure) of the considered conformers, ordered by increasing energy. The supermolecular reference calculations are plotted as solid black line ("Singlepoint").

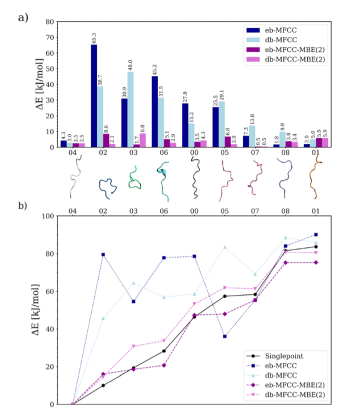


Fig. 7 Comparison of the energy-based and the density-based MFCC and MFCC-MBE(2) schemes for a test set of nine conformers of the (Ala)₁₀ β strand. (a) Absolute errors in the total energy (DFT/BP86/DZP) compared to a single-point calculations for the full polypeptides. (b) Relative energies (DFT/ BP86/DZP) (i.e., energy difference to the lowest-energy structure) of the considered conformers, ordered by increasing energy. The supermolecular reference calculations are plotted as solid black line ("Singlepoint").

for the eb-MFCC and the db-MFCC scheme are even more pronounced than for the turn structures. The inclusion of two-body contributions reduces the errors substantially in both the energy-based and the density-based schemes. Both eb-MFCC-MBE(2) and db-MFCC-MBE(2) result in similar absolute errors, which fall between 0 and 9 kJ mol⁻¹. While neither eb-MFCC nor db-MFCC are able to reproduce the relative energies correctly, both the eb-MFCC-MBE(2) and db-MFCC-MBE(2) show a good agreement of the overall energy patterns.

Across all three structures, only the db-MFCC-MBE(2) scheme is able to provide consistent relative energies for the different polypeptide conformers. In cases with strong intramolecular interactions, such as those present in the α -helical structures, the inclusion of the density-based correction is essential for an accurate description of these interactions. For the turn and βstrand structures, the intermolecular interactions can largely be captured already with the eb-MFCC-MBE(2) scheme.

4 Conclusions

We have combined the eb-MFCC-MBE(2) scheme, which is a simple fragmentation method for proteins that includes a consistent two-body correction, with the density-based many-body **Paper**

expansion previously developed for molecular clusters. The

db-MBE provides a density-based correction to the total energy, which can be calculated from only the electron densities of the considered fragments, i.e., no additional quantum-chemical calculations are required.

For the considered test cases, we could demonstrate that the inclusion of such a density-based correction improved the accuracy considerably, while adding little computational overhead. For idealized polypeptide structures, the db-MFCC-MBE(2) scheme is able to bring down the fragmentation errors to ca. 1 kJ mol⁻¹ for all considered secondary structure elements. For proteins, we are able to reach errors below 1 kJ mol⁻¹ per amino acid. However, these still correspond to substantial errors in the total energies of the proteins, even though the db-MFCC-MBE(2) scheme clearly improves upon the corresponding energy-based scheme.

When considering the relative energies of polypeptides, the db-MFCC-MBE(2) scheme is the only method that is able to consistently provide accurate energies across all considered structural motifs. This underlines the potential of the db-MFCC-MBE(2) scheme as starting point for the parametrization of machine-learning potentials for the use in molecular dynamics simulations of ground and excited states.

While in the present work, we only combined the db-MFCC-MBE(2) scheme with DFT calculations, it is not limited to specific quantum-chemical methods. The scheme presented here is straightforward to generalize to highly accurate wavefunctionbased methods, such as coupled-cluster theory, as previously demonstrated for molecular clusters.⁶⁰

Similarly, our density-based approach is not limited to the MFCC-MBE(2) scheme, but can be combined with any energybased fragmentation method. Therefore, it might be worthwhile to explore the effect of different fragmentation and capping schemes on the fragmentation error.

For further improving the accuracy of the fragmentation method presented here, the combination with a suitable embedding scheme for the fragment calculations seems most promising. Here, we plan to explore both a point-charge embedding as well as the density-based 3-FDE scheme. 81 Finally, the extension of the eb-MFCC-MBE and db-MFCC-MBE schemes to higher orders in the many-body expansion is straightforward.

Author contributions

Johannes R. Vornweg: methodology (lead), software (lead), visualization (lead), writing - original draft (lead), writing review and editing (equal); Toni M. Maier: methodology (supporting), software (supporting), writing - original draft (supporting), writing - review and editing (equal); Christoph R. Jacob: conceptualization (lead), methodology (supporting), software (supporting), writing – review and editing (equal).

Data availability

Data for this paper, including PDB files of all considered molecular structures, Jupyter notebooks for data analysis

(including all raw data) and for generating all figures, as well as PyADF input scripts to perform the eb-MFCC, db-MFCC, eb-MFCC-MBE(2) and db-MFCC-MBE(2) calculations, are available in Zenodo at https://doi.org/10.5281/zenodo.14884143 (ref. 111).

Conflicts of interest

There are no conflicts to declare.

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