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Unlocking the predictive power of quantum-inspired representations for intermolecular properties in machine learning†

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The quest for accurate and efficient Machine Learning (ML) models to predict complex molecular properties has driven the development of new quantum-inspired representations (QIR). This study introduces MODA (Molecular Orbital Decomposition and Aggregation), a novel QIR-class descriptor with enhanced predictive capabilities. By incorporating wave-function information, MODA is able to capture electronic structure intricacies, providing deeper chemical insight and improving performance in unsupervised and supervised learning tasks. Specially designed to be separable, the multi-moiety regularization technique unlocks the predictive power of MODA for both intra- and intermolecular properties, making it the first QIR-class descriptor capable of such distinction. We demonstrate that MODA shows the best performance for intermolecular magnetic exchange coupling (J_{AB}) predictions among the descriptors tested herein. By offering a versatile solution to address both intra- and intermolecular properties, MODA showcases the potential of quantum-inspired descriptors to improve the predictive capabilities of ML-based methods in computational chemistry and materials discovery.

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Introduction

Machine Learning (ML) is having a tremendous impact in Quantum Chemistry (QC).^{1–6} Several research fields are benefiting from new computational strategies combining QC and ML, such as molecular electronics,^{7,8} excited states,^{9–11} low cost discovery of materials,^{12,13} or catalysis.¹⁴ Generally, ML models require data to be transformed into a fixed-size *representation*, usually in the form of a vector, where each element represents a specific attribute or feature. In chemistry, the construction of these elements (typically called *descriptors*) is particularly challenging due to the diversity and complexity of

chemical systems and their interactions. Unlike other applications, molecules do not possess an intrinsic order, structure or size, which makes it difficult to define a universal representation. In the last decade, significant efforts have been made to develop reliable descriptors.¹⁵ These can be classified in three different categories. The first one are cheminformatics descriptors, based on either string fingerprints^{16,17} or on descriptive properties that are easily obtainable by *a priori* knowledge, such as the number of aromatic rings or the molecular size.^{18,19} The second category comprises descriptors based on three-dimensional structural information, usually supplemented with parameters inherited from classical mechanics (*e.g.*, electrostatic potentials, or local density overlaps). In essence, this category does not consider the principles of quantum mechanics and, thus, the descriptors belonging to this category can be referred to as classical-informed representations (CIR). Examples include SOAP²⁰ (Smooth Overlap of Atomic Positions), SLATM²¹ (Spectrum of London and Axilrod–Teller–Muto), many-body interaction descriptor,²² MBTR²³ (Many-Body Tensor Representation), BoB²⁴ (Bag of Bonds) or ACSF²⁵ (Atom-Centered Symmetry Functions). Finally, the third category are descriptors that utilize principles of quantum mechanics to represent the molecular systems and, hence, can be referred to as quantum-informed representations (QIR), with SPA^HM²⁶ (Spectrum of Approximated Hamiltonian Matrices Representation), FJK²⁷ (Fock–Coulomb-Exchange) and MAOC²⁸ (Matrix of Orthogonalized Atomic Orbital Coefficients) being the few existing ones.

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† Electronic supplementary information (ESI) available: (1) Assessment of the effect of the basis set in MODA intermolecular components. (2) Decoupling formalism used to separate intra- and intermolecular components of BoB and SOAP; methodology based on spectral clustering for automatic detection of moieties in multi-moiety systems. (3) Detailed description of the TTTA dataset; two-sample Kolmogorov–Smirnov test for HT-300K and HT-250K AIMD simulations. (4) Results of agglomerative clustering for BoB, SOAP, and MODA with a varying number of clusters. (5) Formalism and strategy to calculate natural orbitals from the density matrix, including an extension for approximating multi-moiety NOs from linear combinations of NOs; representation of the SONOs for TTTA, THIL, and PHYL included in the MODA representation. (6) Analysis of MODA and BoB results for intra- and intermolecular decoupling. (7) Description of the leave-p-groups-out cross-validation strategy used in the KRR predictions. (8) Notes on the performance of descriptors. See DOI: <https://doi.org/10.1039/d3dd00187c>



The main advantage of QIR over CIR descriptors is that the former can explicitly encode the electronic state of a system, including its electronic structure, charge and spin multiplicity. These attributes are necessary to discriminate, *e.g.*, radicals from closed-shell molecules, or neutral from charged species, particularly when the molecular geometry is not significantly affected (and thus, CIR does not significantly change). Although in their early stage of development, QIR descriptors are believed to have the potential to enhance the prediction of properties rooted in the electronic structure of a system. However, a common problem for both CIR and QIR approaches when combined with ML models is the prediction of intermolecular properties.^{29,30} The reason is that, for the sake of better sensitivity, the majority of the descriptors emphasize the magnitude of short-range interactions to better capture the local atomic environment. As a result, these representations can underperform in cases where medium- or long-range (possibly intermolecular) interactions are crucial.^{15,31} This issue can be addressed by decoupling intra- and intermolecular interactions in the representation, and simply ignoring the former in the construction of the descriptor.^{29,32} Although this approach has proven to be successful, neglecting the intramolecular components might be a rather severe simplification, since they can be correlated to the intermolecular property to some extent. For this reason, we here present a mathematical formalism able to accommodate fine-tuned contributions of intra- and intermolecular components in kernel-based ML models.

In general, the possibility to decouple atomic interactions depends on the strategy used to encode molecular information. For instance, BoB is based on 2-body Coulomb potentials and, hence, it can be decoupled using some distance or connectivity criterion. Contrarily, SPA^HM and MAOC, being based on an eigen-decomposition of the Hamiltonian, are not compatible with such an approach, as the resulting eigen-states are related to many-body interactions, potentially gathering contributions from atoms in different moieties. The development of methods that combine the advantages of QIR descriptors for fine-grained representations and the separability of some CIR are sought, as would enable accurate predictions of intermolecular properties while maintaining the benefits of QIR approaches. The relevance of such methods is thus clear, as it would significantly extend the capabilities of ML models in predicting complex intermolecular properties. Herein, we introduce the Molecular Orbital Decomposition and Aggregation (MODA) as a new QIR descriptor, the first of its kind that allows decoupling strategies. As in SPA^HM and MAOC, MODA does not require the calculation of self-consistent field (SCF) solutions. Instead, MODA representation can be constructed using well-established “guess” Hamiltonians,³³ such as the Superposition of Atomic Densities (SAD),³⁴ the Superposition of Atomic Potentials (SAP),³⁵ or the extended Huckel method³⁶ (EH), which are typically starting points in quantum chemistry. The use of these “guess” Hamiltonians provides a computationally-light, yet powerful and simple framework to develop QIR descriptors.

One adequate platform to test and develop new QIR descriptors and also evaluate the possibility to separate intra- and intermolecular components is molecular magnetism. In

particular, we have focused on the evaluation of magnetic exchange couplings, J_{AB} , which quantify the strength and character of the spin-spin interactions between two, *A* and *B*, spin carrying moieties. As it has been shown for datasets comprising di-copper complexes described by BoB, SOAP and MBTR, among others, the prediction of J_{AB} with non-linear regression models can be challenging.³⁷ Moreover, magnetic interactions often occur through-space between independent molecular units and, thus, J_{AB} becomes an intermolecular property. In the following sections, (a) we present the formalism to derive MODA and the strategy to decouple it in intra- and intermolecular components, (b) we discuss the modifications required to kernel-based ML models to accommodate the decoupled representations and, (c) we assess the performance of MODA and other descriptors at predicting magnetic exchange couplings, J_{AB} , as a representative example of an intermolecular property.

In this work, we implement decoupled versions of BoB and SOAP, and we show its advantages over the standard versions when intermolecular properties are targeted. Additionally, we prove that MODA outperforms the decoupled versions of BoB and SOAP in standard supervised regression methods and unsupervised classification models, such as Kernel Ridge Regression³⁸ (KRR) and Agglomerative Clustering³⁹ (AC). Overall, we will demonstrate that MODA is the best representation, among the descriptors here tested, to tackle the prediction of intermolecular J_{AB} interactions, while still being suitable for intramolecular ones.

Methodology

Mathematical formalism

The computation of MODA starts from the atomic positions, which are the only required input (see Fig. 1, step 1). Subsequently, molecular orbitals, the core mathematical object to assemble the representation, are calculated, and expanded as a linear combination of atomic orbitals (see Fig. 1, step 2):

$$|\psi_k\rangle = \sum_i c_{ik} |\chi_i\rangle \quad (1)$$

where c_{ik} are the coefficients describing the contribution of each atomic orbital, $|\chi_i\rangle$, to the molecular orbital, $|\psi_k\rangle$. From this set of coefficients, we formulate the density matrix (D) as a sum of partial density matrices associated to each molecular orbital (D_k).

$$D = \sum_k n_k |c_k\rangle \langle c_k| = \sum_k D_k \quad (2)$$

where n_k corresponds to the orbital occupation. We can measure the interaction strength between pairs of atoms in the system using the density matrix of the molecule (eqn (2)) by means of the Mayer's definition of bond order.⁴⁰ First, the matrix $P = DS$ is constructed, where the overlap matrix (S) is included to account for the non-orthogonality of the atomic orbital basis set. Subsequently, the bond order between two atoms (\mathcal{B}_{AB}) is determined by the trace of the product of P_{AB} and P_{BA} blocks, which contain the rows and columns of P associated





Fig. 1 Scheme of the workflow to compute the features of MODA. The figure shows the process that produces MODA features from the molecular structure to the final representation.

with the atomic orbitals centered at atoms A and B . Formally, the Mayer's bond order is defined as

$$\mathcal{B}_{AB} = \sum_{i \in A} \sum_{j \in B} p_{ij} p_{ji} = \text{Tr}(P_{AB} P_{BA}) \quad (3)$$

where i and j run over all the basis functions centered at atoms A and B , respectively, and p_{ij} are the elements of the matrix P . Exploiting the decomposition of the density matrix into orbital contributions, as stated in eqn (2), the bond order can be expressed as a sum of partial bond orders (\mathcal{B}_{AB}^k) each corresponding to the contribution of a specific molecular orbital, $|\psi_k\rangle$, to \mathcal{B}_{AB} (see Fig. 1, steps 3 & 4).

$$\mathcal{B}_{AB} = \sum_k \text{Tr}(P_{AB}^k P_{BA}^k) = \sum_k \mathcal{B}_{AB}^k \quad (4)$$

The contributions to the partial bond can be decomposed even further. Specifically, we can examine the contributions stemming from the different atomic orbitals of $|\psi_k\rangle$. To this end, we introduce an orbital-specific partial bond order contribution, here represented as $\mathcal{B}_{AB}^{k,nn'l}$. In this notation n and n' correspond to the principal quantum number of the basis functions of atoms A or B , while l corresponds to the azimuthal quantum number. Building upon that, the centerpiece strategy

to generate MODA components consists of the aggregation of such orbital-specific partial bond orders in many-body batches to guarantee the translational, permutational and rotational invariance of the representation. Mathematically, MODA representation corresponds to the set $\{\rho_{nn'l}^{Z_A Z_B}\}$, where each element is defined as (see step 5 in Fig. 1):

$$\rho_{nn'l}^{Z_A Z_B} = \sum_k \sum_{A \in Z_A} \sum_{B \in Z_B} \mathcal{B}_{AB}^{k,nn'l} \quad (5)$$

here A and B refer to all possible atoms in a given molecule with atomic numbers Z_A and Z_B , respectively. That is, each term in MODA aggregates the partial bond orders of all A - B interactions coming from specific atomic orbitals univocally identified by n , n' and l quantum numbers, allowing all possible values of the magnetic quantum number, m (see step 6 in Fig. 1). Our choice of using one azimuthal quantum number while incorporating two principal quantum numbers is inspired by the level of nuance present in the standard SOAP representation. Nevertheless, alternative versions of MODA could delve deeper into distinguishing components by incorporating another quantum number, l' . This would allow to differentiate, for example, σ - σ from σ - π interactions. However, this extra level of nuance would also come at the cost of enlarging the representation size.



Intra-/intermolecular decoupling strategy in MODA

As MODA batches result from the aggregation of pairwise interactions, it can be adapted to decouple intra- and intermolecular interactions. This capability can be incorporated directly into our formalism through a minor modification of eqn (5). For instance, consider δ_{AB} as a parameter that takes a value of 1 when atoms A and B belong to the same molecule, and 0 otherwise. Accordingly, MODA components ascribed to intra- and intermolecular interactions (with super-index I and i , respectively) can be defined as

$$\begin{aligned}\rho_{nn'I}^{Z_A Z_B, I} &= \sum_{\forall k} \sum_{A \in Z_A} \sum_{B \in Z_B} \mathcal{B}_{AB}^{k, nn'I} \delta_{AB} \\ \rho_{nn'I}^{Z_A Z_B, i} &= \sum_{\forall k} \sum_{A \in Z_A} \sum_{B \in Z_B} \mathcal{B}_{AB}^{k, nn'I} (1 - \delta_{AB})\end{aligned}\quad (6)$$

Although eqn (5) and (6) consider the contribution of all molecular orbitals (see the range of k in the outermost summation), we can tune MODA components to include only a S subset of orbitals, denoted as $k \in S$. This is beneficial to avoid the computation of bond orders between orbitals with negligible contributions (e.g., core orbitals), or when the property under examination arises primarily from a specific group of molecular orbitals. For instance, here we have selected the singly-occupied natural orbitals (SONOs) to interpret our property of interest, J_{AB} . Accordingly, the described methodology can be applied to any set of orbitals, and allows the flexible selection of both the level of theory and the basis set, which should be considered as hyperparameters of the MODA representation.

In line with other works in this field,²⁶ our implementation of MODA employs the SAD “guess”, which is a good compromise between computational efficiency and quality of the approximate electronic structure. Note that, resembling the impact of basis sets in quantum chemistry computations, larger basis sets lead to better MODA representations and, thus, better predictions (see Section 1 in the ESI†).

Multi-moiety decoupling

Many ML models, such as Kernel Ridge Regression (KRR) or Agglomerative Clustering (AC), utilize the kernel trick^{41,42} to map the features space into samples space, allowing to use non-linear metrics in practice. The radial basis function (RBF) is a common kernel of choice, which can be defined as

$$\kappa(\rho_1, \rho_2; \gamma) = e^{-\gamma \|\rho_1 - \rho_2\|^2} \quad (7)$$

where ρ_1 and ρ_2 are vector representations of two data samples computed by an arbitrary descriptor, γ is a hyperparameter that determines the decay of the RBF kernel, and $\|\cdot\|$ is the euclidean distance. When the representation ρ can be split in intra-molecular (ρ^I) and intermolecular features (ρ^i), the euclidean distance can be arranged as:

$$\|\rho_1 - \rho_2\|^2 = \|\rho_1^I - \rho_2^I\|^2 + \|\rho_1^i - \rho_2^i\|^2 \quad (8)$$

Consequently, the RBF kernel can be expressed as the product of intra- and intermolecular kernels, where γ is

replaced by two hyperparameters (γ^I, γ^i) that control the width of intra- and intermolecular components, respectively. The definition of the RBF kernel (eqn (7)) can then be combined with the former splitting (eqn (8)) as

$$\begin{aligned}\kappa(\rho_1, \rho_2) &= e^{-\gamma^I \|\rho_1^I - \rho_2^I\|^2} e^{-\gamma^i \|\rho_1^i - \rho_2^i\|^2} \\ &= \kappa(\rho_1, \rho_2; \gamma^I) \kappa(\rho_1, \rho_2; \gamma^i)\end{aligned}\quad (9)$$

The use of separate hyperparameters for intra- and intermolecular components provides a more nuanced approach to model molecular interactions of multi-moiety systems, as the ML model can calibrate the effect of γ^I and γ^i separately. Moreover, the user can aid in the interpretation of the model's predictions by simply inspecting $\gamma^I/(\gamma^I + \gamma^i)$. However, it is important to consider the trade-off between the added flexibility and the increased difficulty in training the model. As the number of hyperparameters increases, so does the effort required to properly fit the model to the data. Additionally, having a larger number of hyperparameters also increases the risk of overfitting, leading to poor generalization performance on unseen data. Therefore, it is crucial to balance the benefits of having more fine-tuned control of the interactions against the increased complexity and the risk of overfitting. We consider the former intra-/intermolecular splitting of the kernel as a reasonable balance between model complexity and interpretability. In addition, note that the here-derived formalism is not specific of the RBF kernel, instead, it can be adapted to a Laplacian kernel or cosine-similarity-based kernels, among other choices.

Finally, it is crucial to note that the formalism derived herein requires the detection of the intra- vs. intermolecular origin of the contributions. Consequently, the algorithm assigned to this task should be informed of the nature of the interaction occurring between each pair of atoms. Given that datasets customarily encompass hundreds or even thousands of data samples, it is crucial that this process is automated in practice to facilitate its efficient application. To meet this requirement, we have devised an algorithm anchored in spectral clustering elaborated in detail in the Section 2 in the ESI.†

Dataset

Next, we introduce all the datasets chosen to illustrate the performance of MODA, prior to present all results and discuss them. Three different datasets have been used in this study to investigate the connection between structural representation and J_{AB} . All three focus on purely organic radicals, since they undergo massive variations in J_{AB} as a result of subtle structural changes.⁴³ The first two datasets are computationally tailored to explore changes along a specific internal coordinate of the system to reduce the structural complexity, providing a simple and chemically intuitive platform to expose the differences between CIR and QIR descriptors. The third dataset comprises a challenging and experimentally realized scenario that includes changes in all structural degrees of freedom in order to test the performance of the evaluated descriptors under more complex conditions.



The first *ad hoc* dataset consists of 60 different conformations of phenalenyl (PHYL, see Fig. 2a) radical pairs, arranged at a fixed inter-planar distance of 3.50 Å and varying the twist in the stacked dimer, θ , from fully eclipsed to staggered configurations ($\theta = 0^\circ$ and 60° , respectively). This dataset provides insight into the influence of relative orientation and molecular orbital symmetry on intermolecular J_{AB} couplings.⁴⁴ The second *ad hoc* dataset, containing 180 different conformations of Thiele's diradical⁴⁵ (THIL, see Fig. 2b), spans the central phenyl ring's rotational angle from $\theta = 0^\circ$ to 180° (see θ in Fig. 2b). Notice that the substantial bulkiness of the terminal phenyl groups induces an out-of-plane twist to alleviate steric hindrance. As a result, the symmetry of the molecule in the planar conformation diminishes from D_{2h} to either D_2 or C_{2h} . Our dataset is based on the rotation of the central phenyl ring of the C_{2h} conformer along the axis connecting the two $-\dot{\text{C}}\text{Ph}_2$ moieties. Note that this rotation implies a sweeping range from $\theta = 0^\circ$ to 180° without mirror symmetry at 90° (refer to Section 5 in the ESI† for a 3D representation of the THIL diradical). This dataset provides a valuable test case for analyzing variations of intramolecular J_{AB} as a result of conformational changes.^{46,47}

The third dataset is made of dimers of 1,3,5-trithia-2,4,6-triazapentalenyl (TTTA see Fig. 2c). In the solid state, TTTA molecules experimentally arrange forming labile 1D π -stacks⁴⁸

(see Fig. 2d). At low temperatures, these remain as alternated dimers but, at higher temperatures, a Pair-Exchange Dynamics (PED) process is triggered, in which the inter-planar distance between a given pair of TTTA molecules oscillates between *ca.* 3.25 Å and *ca.* 4.5 Å.⁴⁹ Such subtle structural changes lead to massive time- and temperature-dependent variations in the J_{AB} interaction between pairs of π -stacked TTTA units,⁵⁰ ranging from weakly ferromagnetic (FM, *ca.* 50 cm^{-1}) to strongly anti-ferromagnetic (AFM, *ca.* -5000 cm^{-1}), as shown in Fig. 2e.

Following the approach of previous works for sample mining,^{7,30,32,51} we obtained TTTA dimers from *ab initio* molecular dynamics (AIMD) simulations of the high-temperature (HT) phase, carried out at two different temperature conditions, 300 K (HT-300K) and 250 K (HT-250K), in order to explore different regions of the thermally available configurational space. Each AIMD simulation had an independent 10 ps run, yielding a variety of structures. The data from the HT-300K phase, detailed in a previous publication⁴⁹ by some of the present authors, included 30 000 configurations selected from 3 different dimers of the crystal structure (10 000 of each one). These configurations have been used to train and test the ML models, offering a diverse training set that encapsulates a wide range of structural variations. Conversely, the HT-250K phase data is a new addition, generated specifically for this study. We

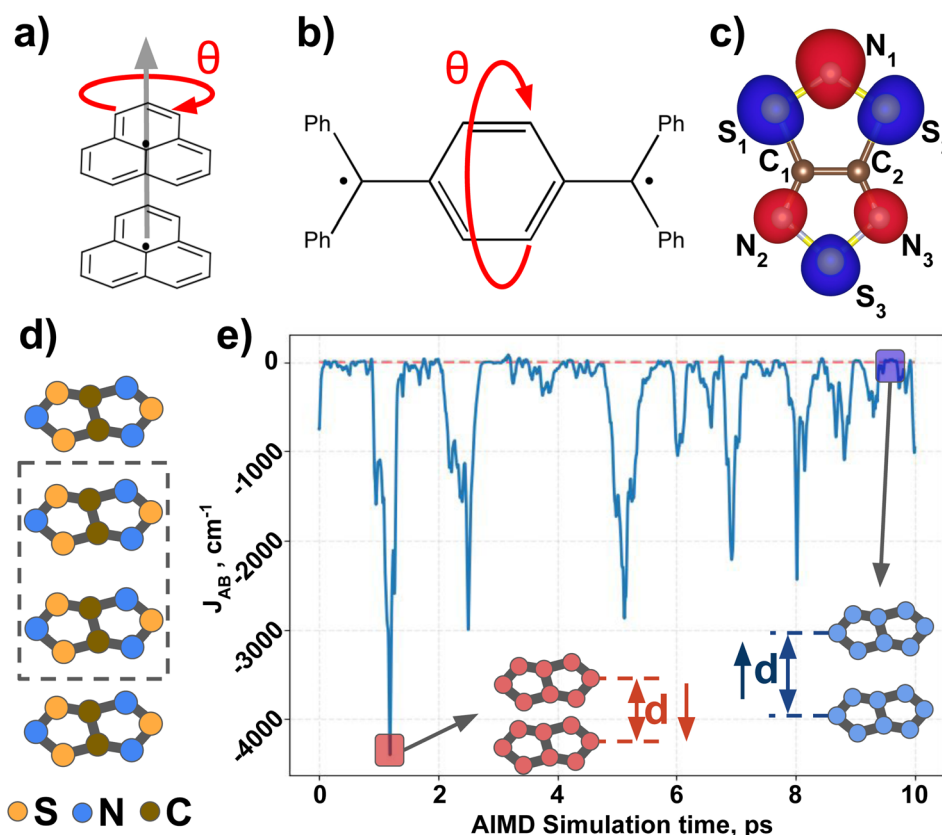


Fig. 2 (a) π -Stacked pair of phenalenyl radicals (PHYL), (b) Thiele's diradical (THIL) and (c) chemical representation overlapped with the SONO of TTTA. Red arrows indicate the rotational internal coordinate (θ) explored to obtain all geometries of PHYL and THIL. (d) π -Stack of TTTA radicals, with a dimer highlighted. (e) Time resolved evolution of J_{AB} of two TTTA radicals forming a dimer along a time interval of 10 ps of AIMD simulation. Insets highlight two regions with extremely different J_{AB} values: -4300 cm^{-1} (red) and $+20 \text{ cm}^{-1}$ (blue).



extracted 20 000 structures from this phase, following the same selection strategy. The HT-250K structures have served as a separate dataset for evaluating the extrapolation capabilities of the models, ensuring that our ML models can make accurate predictions on unseen data spanning different regions of the configurational space. A detailed analysis of the TTTA dataset can be found in Section 3 of the ESI†

Results and discussion

The discussion of the results is organized into five subsections. In the first part, we evaluate the capabilities of CIR and QIR descriptors in unsupervised learning tasks involving J_{AB} predictions by means of Agglomerative Clustering (AC). The second section demonstrates the benefits of separating intra- from intermolecular degrees of freedom in descriptors when predicting intermolecular J_{AB} values for both CIR and QIR-type representations. In the third section, we highlight the key qualitative differences between CIR and QIR-type descriptors using the TTTA dataset. Complementing the former, the fourth section validates quantitatively the observations discussed in the previous section using variance-covariance and mutual information analyses. Lastly, in the fifth section, we provide additional evidence supporting MODA's superior performance in TTTA dataset by presenting our findings from KRR predictions. Throughout the discussion, we compare structure-average SOAP and MODA as the main representative examples of CIR and QIR-class descriptors, respectively, that enables the decoupling of intra- and intermolecular components (see similar analyses for BoB in Sections 4 and 6 of the ESI†). As mentioned in previous sections, MODA describes a specific subset of molecular orbitals. In this case, the MODA representation of the systems under study is constructed using their respective SONOs, since these MOs play a crucial role in determining the J_{AB} values⁵² (see more information in Section 5 of the ESI†).

CIR vs. QIR performance on *ad hoc* datasets

PHYL and THIL datasets focus on an intermolecular J_{AB} with fixed intra-molecular components, and a fully intramolecular J_{AB} , respectively. As introduced in the dataset section, both datasets have been specially designed to simplify the structural variability to a single internal coordinate (a rotational angle, θ). DFT calculations have been performed along this coordinate to retrieve θ vs. J_{AB} curves, and the clusters resulting from the AC model are projected onto these curves (see Fig. 3 for PHYL and Section 4 of the ESI† for THIL).

The J_{AB} curve of PHYL spans a wide range of values from strongly AFM couplings in eclipsed and staggered conformations (with D_{3h} and D_{3d} point group symmetries, respectively) to slightly FM couplings at $\theta = 30^\circ$ (S_6 point group), and displays symmetry around $\theta = 30^\circ$ (see Fig. 3a). The mirror-symmetric profile of J_{AB} around $\theta = 30^\circ$ (see Fig. 3a) is at odds with the point group symmetry associated to the PHYL's conformers along θ ($D_{3h} \rightarrow S_6 \rightarrow D_{3d}$). The reason for such discrepancy can be grasped by the electronic structure of the SONOs of the PHYL



Fig. 3 J_{AB} profile of PHYL conformers from eclipsed ($\theta = 0^\circ$) to staggered ($\theta = 60^\circ$) geometries. The color code indicates the clustering produced by (a) SOAP and (b) MODA descriptors. The insets of (a) indicate the evolution of the point group symmetry at specific values of θ associated with the geometry of the dimer. Analogously, the insets in (b) indicate the point group symmetry associated with the SONOs. The bottom part of the figure illustrates the SONOs at $\theta = 0^\circ$ and $\theta = 60^\circ$, in which the highlighted hydrogen shows that the structure is different, while the SONO remains unaltered.

dimer. Both SONOs at the eclipsed and staggered conformations belong to D_{3h} point group, while the $\theta = 30^\circ$ conformer belongs to the D_{3d} point group in this case (see SONOs in Fig. 3b). Thus, the SONOs of the PHYL conformers from $\theta = 0^\circ$ to 30° ($D_{3h} \rightarrow D_{3d}$) are mirror-symmetric to the conformers from $\theta = 30^\circ$ to 60° ($D_{3d} \rightarrow D_{3h}$), just as the J_{AB} curve along θ . That is, the relevant symmetry to interpret the J_{AB} evolution is not the symmetry associated to the geometry, but the one emerging from the electronic structure.⁵³ Consistent with this observation, the clustering patterns of SOAP (which is



exclusively based on structural information) fail to capture the symmetry around $\theta = 30^\circ$, and instead follow the clustering pattern arising from structural symmetries (Fig. 3a). In contrast, MODA, being a QIR descriptor, produces a mirror-symmetric clustering pattern around $\theta = 30^\circ$, in agreement with the molecular orbital symmetry associated to J_{AB} (Fig. 3b).

It is worth noting that the use of structure-average SOAP vectors might not fully leverage the potential of the descriptor. This is particularly relevant given that the performance of clustering techniques can be significantly impacted by minor distortions in the distance between samples. To substantiate our comparison between SOAP and MODA further, we expanded our investigation to include the use of a more general local SOAP representation, employing the Regularized Entropy Match⁵⁴ (REMatch) together with RBF kernel to evaluate AC performance from the local SOAP representation. This exploration consistently validated our initial observations regarding the structure-average version of the descriptor, as detailed in Section 4.2 of the ESI†

To further extend the analysis of the capabilities of SOAP and MODA, we assessed their performance using the THIL dataset, which features a molecular diradical instead of a pair of radicals. Here, J_{AB} is geometrically controlled by the through-bond conjugation of both $-\dot{\text{C}}\text{Ph}_2$ groups to the central phenyl group guided by the angle: J_{AB} is largely AFM when the central phenyl group remains in-plane ($\theta = 0^\circ$ and $\theta = 180^\circ$), and it drops to zero as the phenyl moiety approaches an orthogonal conformation ($\theta = 90^\circ$). Upon examining the electronic structure of THIL no special symmetry arising from the SONOs can be anticipated and, thus, the atomic disposition and electronic structure of the SONOs are bijectively related (see section 5.3 in the ESI†), which suggests that CIR descriptors can well describe J_{AB} in the THIL dataset. Agglomerative clustering results further confirm this observation, showing small variation among SOAP and MODA descriptors with no significant impact on the AC performance (elaborated in detail in Section 4 of the ESI†). This scenario arises when the system, like THIL, is well-characterized by geometry, rendering CIR and QIR-based descriptors, such as MODA, comparably efficient.

Consequently, our analyses underline MODA's superior performance over CIR methods like SOAP and BoB (as further discussed in Section 4 of the ESI†) in scenarios that present challenges to geometry-based descriptors, as evidenced in the PHYL results. This enhanced performance is due to the QIR's inclusion of wave-function information, capturing crucial, otherwise overlooked, electronic structure elements. Nonetheless, in scenarios where the system is properly defined by geometric descriptors, such as in the THIL dataset, both CIR and QIR-based descriptors can achieve similarly proficient results.

Decoupling intra- and intermolecular features in SOAP

In the previous subsection, we focused on cases with controlled *ad hoc* structural changes, in order to illustrate some fundamental differences between CIR and QIR-type descriptors. However, a more appealing challenge for a ML model is the prediction of intermolecular properties in conditions where the

intramolecular degrees of freedom of each moiety are not artificially constrained. To address this scenario, we turn our attention to the TTTA dimers dataset, where the samples originate from AIMD simulations and, thus, the motion of atoms is uncontrolled and conditioned by thermal fluctuations. As demonstrated below, this is the perfect platform to evaluate the importance of decoupling the components of descriptors. Our analysis initially focuses on structure-average SOAP, as a formerly validated choice in PHYL by Agglomerative Clustering experiments, and then extends to MODA.

The advantages of decoupling intra- and intermolecular interactions become clear when comparing the target J_{AB} with the intra- and intermolecular components of SOAP separately (see Fig. 4). Intramolecular features span a larger range of values, showing the tendency of CIR descriptors to emphasize short-range interactions, but these features show no correlation whatsoever with J_{AB} (see Fig. 4a and b). Thus, even if some degree of correlation exists between intermolecular components and J_{AB} (see Fig. 4b and c), the strong and noisy intramolecular components can make the standard (*i.e.*, not decoupled) SOAP struggle when capturing the evolution of J_{AB} values. Fine ML predictions are still possible, since the intramolecular components might still be correlated with intermolecular ones to some extent (*e.g.*, the variation of the bond lengths when forming a dimer) and hence to J_{AB} . However, such predictive models will primarily memorize data rather than generalize the trends governing the descriptor-to-target mapping, resulting in overfitted ML models that struggle to interpolate and extrapolate beyond the available data. Analogously, the result obtained for SOAP regarding the decoupling of intra/inter features is consistent with the observations for BoB and MODA (see Section 6 of the ESI†).

Performance of MODA and SOAP in the TTTA dataset

Having established (i) the importance of having a QIR descriptor for the prediction of J_{AB} values, and (ii) the advantages of the decoupling strategy, we will now discuss the role of both elements in the case of the TTTA dataset. The SONOs of TTTA have dominant S and N contributions, and negligible contribution from its C atoms (Fig. 2c). Accordingly, MODA components related to C–C interactions are negligible in comparison to the components associated with S–S or N–N interactions (see Fig. 5a). Contrarily, SOAP overemphasizes C–C interactions, thereby rendering them comparable to N–N and S–S components across most of the examined range of Fig. 4. These are the three interaction types that exhibit the highest variance, and whose curves jointly vary along the AIMD simulation time (see highlighted red-most curves in Fig. 4c).

Moreover, MODA's features exhibit a stronger correlation with J_{AB} compared to the intermolecular SOAP components discussed above. This correlation becomes apparent when examining the descriptor response ($\Delta\rho$) to changes in J_{AB} over the AIMD simulation time (t). For instance, when TTTA dimers undergo abrupt structural changes that shift their associated J_{AB} from paramagnetic to strongly AFM regimes (large $\Delta_{12}J$, t_1 vs. t_2 in Fig. 5b), MODA accurately captures these changes (large $\Delta_{12}\rho_{\text{MODA}}$,



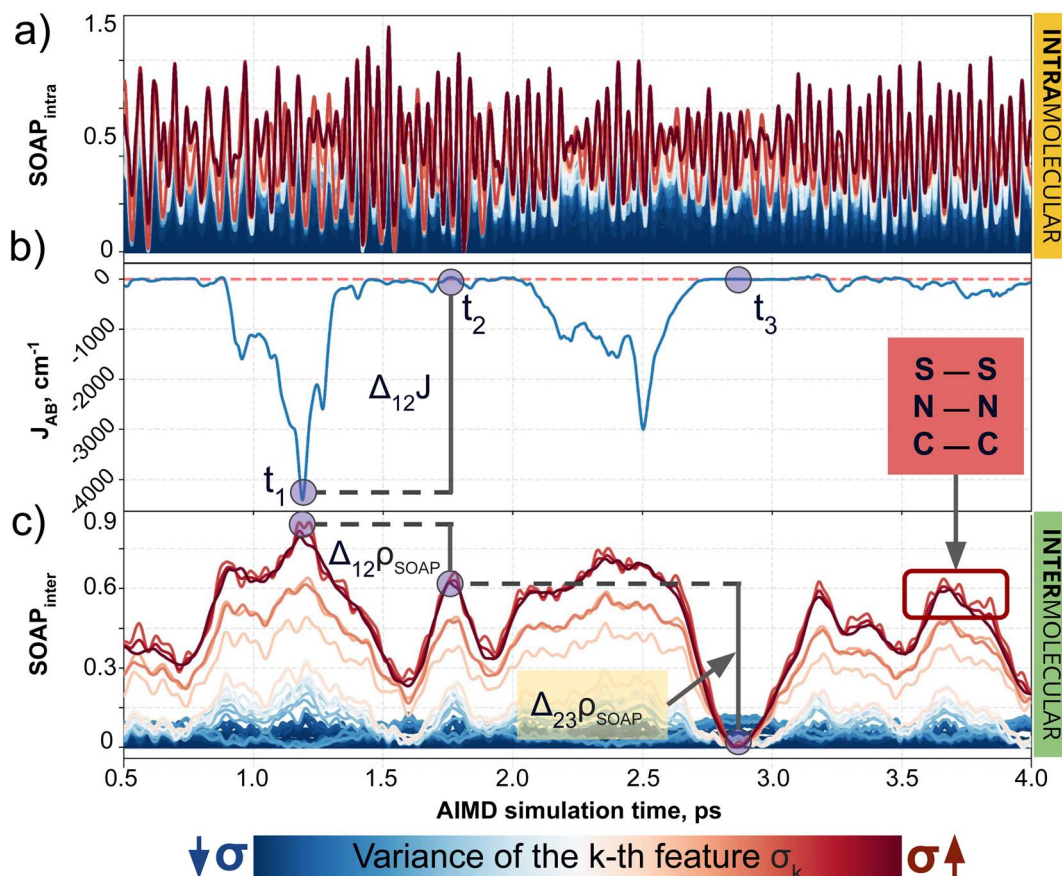


Fig. 4 Evolution of (a) intra- and (c) intermolecular SOAP features (colored by variance value) of a TTTA dimer along the structures explored during the 0.5–4.0 ps time interval of AIMD, as well as (b) the associated time-resolved evolution of J_{AB} in this interval. The blue circles in (b) and (c) indicate three regions with specific values of J_{AB} , where one can compare the change associated with J_{AB} and SOAP.

Fig. 5a). Conversely, SOAP fails to accurately represent these variations (small $\Delta_{12\rho_{\text{SOAP}}}$, Fig. 4c), causing completely opposite structures to appear incorrectly similar. In the same vein, J_{AB} values at $t = t_2$ and $t = t_3$ are almost identical in both cases. In this situation, MODA assigns a similar response to both representations, while SOAP indicates an abrupt change ($\Delta_{23\rho_{\text{SOAP}}}$), even larger than the change associated with the extremely contrasting regimes at $t = t_1$ and $t = t_2$. The primary explanation for this discrepancy is illustrated in Fig. 5d and e, which displays the structure and SONOs of the TTTA dimer at $t = t_2$ and $t = t_3$, see d_{AB} in the figure. Accordingly, the TTTA moieties at $t = t_2$ are notably closer than at $t = t_3$. Nevertheless, the overlap between the SONOs is nearly zero in both cases (compare to the shared isosurface between TTTA SONOs of $t = t_1$ in Fig. 5c), resulting in a similar J_{AB} . This situation is particularly challenging for SOAP, which fails to capture these subtleties due to its lack of electronic structure information. As a result, MODA's ability to capture electronic structure information translates into a higher correlation with J_{AB} than that exhibited by SOAP in this specially critical ill-defined situation for CIR descriptors.

Mutual information and covariance analyses

Our qualitative comparison between descriptors can be complemented using the point-wise Global Feature Reconstruction

Error, $\text{GFRE}^{(t)}(F, F')$. This method involves reconstructing the features of a given descriptor (F') using another one (F), in order to assess the presence/absence of mutual information.⁵⁵ In this case we have assessed how intermolecular components of SOAP can reconstruct MODA, and *vice versa*. For completeness, we have used two flavors of MODA: the one we have used along this section, using only the SONOs ($\text{MODA}_{\text{SONOs}}$), and a complete one in which all the occupied MOs are considered (referred to as MODA_{occ}). The large value around $t = t_3$ in both $\text{GFRE}^{(t)}$ spectra indicates that neither descriptor can reproduce the other, meaning that $\text{MODA}_{\text{SONOs}}$ and SOAP carry substantially different information in their representation (see Fig. 6a). This is expected, since SOAP contains structural information, while $\text{MODA}_{\text{SONOs}}$ contains electronic structure information specially devoted to capture J_{AB} changes. Besides $t = t_3$, we can observe that $\text{MODA}_{\text{SONOs}}$ frequently struggles to reconstruct SOAP, while the opposite rarely happens (see Fig. 6a). The overall perspective can be quantified using the global $\text{GFRE}(F, F')$ (see Fig. 6b), resulting in a reconstruction error that is larger when going from $\text{MODA}_{\text{SONOs}}$ to SOAP (0.531) than the other way around (0.029) (see light blue frames in Fig. 6b). Interestingly, when using MODA_{occ} , mutual reconstruction error with SOAP is similar and low. This suggests that MODA, in its different flavors dictated by the choice of molecular





Fig. 5 Evolution of (a) intermolecular MODA features (colored by variance value) of a TTTA dimer along the structures explored during the 0.5–4.0 ps time interval of AIMD, as well as (b) the associated time-resolved evolution of J_{AB} in this interval. The blue circles in (a) and (b) allow for comparison between variations in J_{AB} and variations in the features of MODA. TTTA dimer's SONOs at three representative time steps: (c) $t = t_1$, (d) $t = t_2$, and (e) $t = t_3$.

orbitals, can range from being a specialized (*e.g.*, $\text{MODA}_{\text{SONOs}}$) to a general (*e.g.*, MODA_{occ}) descriptor. On a lesser note, the reconstruction error associated with BoB, irrespective of

during reconstruction or being reconstructed, is substantially higher compared to the rest of the cases (see red-framed values in Fig. 6b). However, the descriptor accomplishing this task



Fig. 6 (a) Point-wise Global Feature Reconstruction Error $\text{GFRE}^{(t)}(F, F')$ during the 0.5–4 ps time interval of AIMD. The upper panel (red curve) shows the error spectrum associated to the reconstruction of $\text{MODA}_{\text{SONOs}}$ (F') using SOAP (F), while the lower panel (blue curve) corresponds to the reconstruction of SOAP features (F') using $\text{MODA}_{\text{SONOs}}$ (F). Dashed black lines indicate three different time regions discussed in the main text. The matrix in (b) indicates the Global Feature Reconstruction Error $\text{GFRE}(F, F')$ using the descriptors in each row (F) to reconstruct the descriptor indicated in each column label (F'). Some matrix elements are highlighted to facilitate the discussion in the text (using color frames).

more successfully is SOAP, emphasizing the shared CIR origin of both representations.

To further support our conclusions, we have performed a variance–covariance analysis of the intermolecular features of BoB, SOAP and MODA (refer to Fig. 7). For MODA, the features bearing a larger variance (that is, those exerting the greatest influence on the overall molecular descriptor, see Section 8 in ESI† for an elaborate discussion) simultaneously possess the highest covariance with J_{AB} (Fig. 7a). In essence, the components with the highest variance are also those exhibiting the strongest linear correlation with J_{AB} . This finding stands in stark contrast to the results obtained for SOAP and BoB (Fig. 7b and c). For instance, in SOAP, the component with the largest impact (σ_1 in Fig. 7b) demonstrates a covariance with J_{AB} that is comparable to the 6th most influential feature (σ_6 in Fig. 7a) of the MODA descriptor. In addition, the covariance of the 20 top most determining features of SOAP and BoB (338.2 and 162.1, respectively) are smaller than in MODA (598.2). However, we acknowledge that the variance–covariance analysis assumes linear relationships, while the features may generally exhibit non-linear connections with target properties. Indeed, while linear relationships are favored for their simplicity and stronger generalization capabilities that illustrates significant differences of the performance of each descriptor in the TTTA dataset, we are aware that variance–covariance analysis can oversimplify the data complexity. To account for non-linearity, we have performed a Kernel Ridge Regression (KRR), further enhancing our understanding of descriptor behavior.

KRR predictions

In order to assess the effect of non-linear relations between data representation and J_{AB} , we have employed KRR together with RBF kernel. The learning curves produced by the evaluation of optimal KRR models are presented in Fig. 8, where SOAP, BoB and MODA descriptors have been used. Within the cross-validation leave-p-groups out scheme (see Section 7 of ESI† for more details), we have used different fractions of the HT-300K data for training, testing, and searching the optimal hyperparameters for γ^l , γ^i , and α (the regularization parameter of

KRR method), while the model performance is evaluated with data samples from the HT-250K AIMD trajectory (validation set). As discussed in the dataset section, the HT-250K trajectory mostly explores the same potential energy surface region compared to HT-300K, but it also includes unseen regions due to the different thermally-available configurational space (see Section 3.1 of the ESI†), which enables to assess the ability of the model in both interpolation and extrapolation tasks.

For all the descriptors, the test curves (dashed lines in Fig. 8) still exhibit a reduction in MAE beyond a training size of 40%. However, the validation curves (solid lines) reach a saturation plateau at 25% regardless of the descriptor in use. This indicates that each model achieves an adequate training size at 25%, and increasing the data size beyond this point only results in KRR models with similar extrapolation and interpolation capabilities. Concerning the accuracy of the different



Fig. 8 KRR learning curves using BoB (red), SOAP (green) or MODA (purple), using the leave p-groups-out strategy. Solid and dashed lines indicates the data source: HT-300K (test) and HT-250K (validation), respectively.



Fig. 7 Variance–covariance plots for intermolecular components of (a) MODA, (b) SOAP and (c) BoB with J_{AB} . The x-axis on each plot (and the color of each bar) corresponds to the variance of the 20 intermolecular features with highest variance, while the y-axis corresponds to the covariance of the component with J_{AB} . Insets show the cumulative covariance of all the plotted features.



descriptors, KRR models based on BoB and SOAP display MAEs around 40 cm^{-1} and 28 cm^{-1} , respectively. In contrast, the KRR model utilizing MODA representation outperforms both SOAP and BoB, registering a substantially lower MAE under 20 cm^{-1} . While the test curve for BoB and SOAP (dashed red and dashed green lines in Fig. 8) reaches the lowest MAE, their respective validation curves (solid red and solid green lines) register substantially higher MAEs in both cases. In contrast, test and validation curves employing MODA (dashed and solid purple lines in Fig. 8, respectively) show a joint evolution along the entire range, which indicates that KRR model trained upon MODA does not present as much overfitting as in BoB and SOAP. These differences reinforce the superior performance of MODA in predicting J_{AB} values in the TTTA dataset, making it a better choice for capturing the underlying relationships in the data and generating more accurate predictions of J_{AB} at a reasonable computational cost (see Section 8 in the ESI†).

Finally, it is pertinent to place the improvement demonstrated in MODA in a proper context. While the overall range of J_{AB} extends approximately from -4000 cm^{-1} to $+100\text{ cm}^{-1}$, a detailed examination reveals that the data points predominantly cluster around 0 cm^{-1} (as detailed in Fig. 3 of the ESI†). The distribution is significantly skewed, with the mean J_{AB} value around 300 cm^{-1} , and a median, which is a more appropriate measure of central tendency for this skewed distribution, approximately at 200 cm^{-1} .

From this standpoint, the MAE of 20 cm^{-1} recorded for MODA corresponds to a 10% error margin with respect to the median, a substantial enhancement compared to the 15% and 21% errors represented by MAEs of 30 cm^{-1} and 42 cm^{-1} for SOAP and BoB, respectively. This improvement is particularly impactful in practical applications. For instance, in molecular magnetism applications, the accurate J_{AB} predictions are crucial, as they serve as intermediate steps in calculating thermodynamic properties like magnetic susceptibility. It is well-established that the prediction of such macroscopic observables from first-principles J_{AB} calculations is highly sensitive to even small variations in J_{AB} , especially in regions nearing 0 cm^{-1} .⁵⁶ Therefore, the enhanced precision of J_{AB} prediction offered by MODA is often of critical importance.

Conclusions

We report the development, implementation and validation of the Molecular Orbital Decomposition and Aggregation (MODA) approach, a novel QIR-class descriptor that encodes the electronic structure of molecules. MODA uses the density matrix of a subset of molecular orbitals to euclidate the contributions of atom pairs as bond orders, and groups these interactions by atom types and classes of atomic orbitals, leading to an invariant representation for molecules. MODA is designed to work in multi-moiety regularization schemes, where the kernel is split into intra- and intermolecular components. We demonstrate that this particular setup results in a better performance in unsupervised and supervised learning tasks targeting intermolecular interactions.

Our study offers a detailed analysis comparing the performance of MODA to some separable CIR descriptors. The examination of pairwise sample-sample maps, along with the application of unsupervised Agglomerative Clustering on *ad hoc* PHYL and THIL datasets, emphasize the importance of capturing the key electronic structure information for an accurate representation of properties rooted in quantum mechanics. Notably, our results show that MODA effectively captures the molecular orbital symmetry, which is crucial for the classification of J_{AB} values, as opposed to other CIR-type representations. The performance of MODA has been further assessed with TTTA dataset, which reinforces the importance of this separation in both CIR and QIR-based representations. Particularly, the variance-covariance analysis and KRR predictions provide additional evidence of MODA's superior performance in capturing relationships with J_{AB} compared to other separable CIR representations.

Overall, MODA, as the first separable QIR-type descriptor, shows potential in enhancing the prediction of molecular properties rooted in quantum chemistry. Based on the specific evaluations presented in this study for J_{AB} values, we believe that MODA will be also applicable to other properties such as HOMO energies or transfer integral predictions.

Computational details

Data mining and J_{AB} evaluation

The AIMD simulation of the HT phase of TTTA at 250 K and 300 K used the same computational protocol employed in earlier works describing this molecule,^{49,50} and other organic radicals.^{57,58} A monoclinic supercell was prepared containing 32 TTTA molecules arranged in 8 stacks of radicals, each of them containing 4 radicals. The AIMD simulation was run for *ca.* 10 ps and a time step of 4 a.u. Vanderbilt ultrasoft pseudopotentials⁵⁹ were employed, together with the PBE functional⁶⁰ within the spin unrestricted formalism and the Grimme D_2 correction,⁶¹ and a Γ -point sampling of the Brillouin zone. The Car-Parrinello⁶² propagation scheme was employed as implemented in CPMD,⁶³ using a fictitious mass for the orbitals of 400 a.u. The simulations were performed in the canonical (or NVT) ensemble using Nosé-Hoover chain thermostats.⁶⁴ All J_{AB} values were evaluated as the difference between the energy of the Broken Symmetry⁶⁵ (E_{BS}) and Triplet (E_T) electronic states assuming the Heisenberg hamiltonian, $\hat{H} = -2J_{AB}\hat{S}_A\hat{S}_B$.⁶⁶ E_{BS} and E_T were computed at the UB3LYP⁶⁷⁻⁶⁹/6-31+G*^{70,71} level as implemented in Gaussian09.⁷² We have employed the two-sample Kolmogorov-Smirnov test⁷³ (as implemented in the *scipy* module in Python) on the statistical distributions of both the centroid-centroid distances between TTTA units, and the J_{AB} values associated with structures extracted from the HT-300K and HT-250 K trajectories, to guarantee that the configurational space sampled at 300 K and 250 K is different (see more details in Section 3.1 of the ESI†). Moreover, PHYL and THIL conformations have been obtained *via* a rigid rotational scan of the angle θ , as described in the dataset section, after a geometry relaxation in the triplet state. We have then calculated J_{AB} values of every configuration following the same level of theory



described above for TTTA, and without imposing structural or orbital symmetry (“NoSymm” keyword).

Descriptors and molecular representations

The decoupled version of BoB, SOAP and MODA descriptors has been obtained through our code implemented in Python and collected in the MLcool module (<https://github.com/GEM2-UB/MODA>). The decoupling scheme applied to BoB (refer to Section 2.1 in the ESI†) is similar to MODA, as both are built from 2-body interactions. However, SOAP, as implemented in many packages, is a 3-body descriptor,⁷⁴ and necessitates a different approach. Therefore, we adopted the approach proposed by Cersonsky *et al.* for SOAP decoupling,⁷⁵ detailed in Section 2.2 in the ESI†. In particular, BoB has been implemented from scratch, while we use the DScRibe⁷⁶ module for local and structure-average versions of SOAP are used in the backend. The version of MODA employed in the manuscript computes the natural orbitals (NOs) and occupation numbers by means of the following two steps: (1) the density matrix is obtained by means of the SAD guess (with guess = “atom”) as implemented in pySCF.⁷⁷ (2) A Lowdin's symmetric diagonalization⁷⁸ is performed to solve the generalized eigenvalues equation⁷⁹ ($DSC = Cn$, see details in Section 5.1 of the ESI†). It is important to note that all the molecules considered in this work are radicals, thus, the SAD guess has been computed in the open-shell spin state. Specifically, for THIL, we have computed the density matrix in the triplet state. In contrast, for PHYL and TTTA (both being pairs of radicals), the density matrix was computed for each monomer separately and subsequently, the dimer electronic structure was approximated as a sum of doublets (see more details in section 5.1.1 of the ESI†). SOAP partial power spectrum vector has been calculated on the atomic positions both locally (average = “off” in DScRibe) and in its structure-average version (average = “outer”). We used a Gaussian-type orbital radical decay function and set the n_{\max} , l_{\max} and r_{cut} hyperparameters to 6, 4, and 7, respectively. Moreover, σ has been varied to specific values (0.1, 1, 2, 3 and 10), with no extra radial scaling weighting of the atomic density to let intermolecular components dominate the SOAP spectrum. MODA representations have been computed using different basis sets (STO-6G, 6-31G, 6-31G*, 6-31+G* and aug-cc-pvdz). We found a significant effect of the basis set on the performance of the model (see Section 1 of the ESI†). Only the results of 6-31+G* basis set are reported in the main text, as it has been found to be the best balance for accuracy and computational cost.

Supervised and unsupervised learning

AC and KRR models have been used as implemented in scikit-learn Python's package.⁸⁰ In both cases the kernel = “pre-computed” option was used to provide our implementation of intra/intermolecular separable RBF kernel version (as described in the methodology section and ESI†) transpiled from Fortran95 and capable of parallel computing. For the local SOAP representation we have employed the Regularized Entropy Match kernel⁵⁴ (REMatch) to evaluate global similarities from local representations. In this case we have experimented with different values of the entropic penalty parameter: $\alpha = 0.01$ (best match),

$\alpha = 1.0$ (intermediate) and $\alpha = 10$ (average-like regime), which constitutes a reasonable choice to sweep the spectrum of regimes that this kernel can work in ref. 81 (see Section 4.1 of the ESI†). Before the RBF kernel is computed, the intra- and intermolecular features have been separately normalized to max = 1 in order to guarantee reasonable optimal values of the hyperparameters. In AC experiments, the whole PHYL and THIL datasets have been used with linkage = “complete”,³⁹ which minimizes the maximum distance between observations of pairs of clusters. A varying number of clusters has been used, ranging from 2 to 10 (the latter is shown in the main text and the rest in Section 4 of the ESI†). We have fitted the KRR model according to a custom implementation of a grid search cross-validation (CV) with leave p-groups-out strategy over γ^l and γ^i hyperparameters of the decoupled RBF kernel, and the ridge regularization term (“alpha” in sklearn implementation). We applied the CV search to TTTA structures of the HT-300K dataset computing R^2 , MAE and RMSE mean values across all the $N!/p!(N-p)!$ CV splits (where N and p stand for total number of groups and number of groups out, respectively), and then used data from the HT-250K phase as our validation set. The grid search for the ridge regularization parameters explored 50 equidistant values in a logarithmic scale ranging from 10^{-11} to 10^{-1} . Similarly, 100 values have been tested for γ^l and γ^i in the interval $(10^{-100}, 10^3)$. The learning curves have been produced by selecting a specific number of total groups and groups out N/p , to get 5% (20/1), 10% (10/1), 20% (5/1), 25% (4/1) and 40% (5/2). GFRE(F,F') and GFRE⁽ⁱ⁾(F,F') feature reconstruction measures have been calculated by means of the scikit-matter⁸² Python module, using only intermolecular features of the descriptors. The regression weights for GFRE⁽ⁱ⁾ were calculated from the samples in the time interval $t \in (4, 10)$ ps and evaluated at $t \in (0.5, 4)$ ps, while a strict 50/50 split have been used in GFRE before random shuffling to decorrelate time-evolution samples.

Data availability

The code and tutorial for the MLcool package, including examples for using BoB, SOAP, and MODA, are available at the GitHub repository: <https://github.com/GEM2-UB/MODA>. A tutorial showing the data analysis of this paper is available in the interactive notebook “MODA_tutorial.ipynb” at <https://github.com/GEM2-UB/MODA>. The complete dataset used for this study, encompassing THIL, PHYL, and TTTA datasets, is accessible at Zenodo: <https://doi.org/10.5281/zenodo.8032717>. Other algorithms and analyses have been uploaded as a part of the ESI†. Both the code and dataset are required to be formally cited in the reference section of this article.

Conflicts of interest

There are no conflicts to declare.

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