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# Explaining Conformational Diversity in Protein Families through Molecular Motions

<sup>3</sup> Valentin Lombard<sup>1</sup>, Sergei Grudinin<sup>2,\*</sup>, and Elodie Laine<sup>1,3,\*</sup>

<sup>4</sup> <sup>1</sup>Sorbonne Université, CNRS, IBPS, Laboratoire de Biologie Computationnelle et Quantitative (LCQB), 75005 Paris,
 <sup>5</sup> France

<sup>6</sup> <sup>2</sup>Université Grenoble Alpes, CNRS, Grenoble INP, LJK, 38000 Grenoble, France

<sup>7</sup> <sup>3</sup>Institut Universitaire de France (IUF)

\* corresponding author(s): Sergei Grudinin (sergei.grudinin@univ-grenoble-alpes.fr), Elodie Laine

9 (elodie.laine@sorbonne-universite.fr)

### 10 ABSTRACT

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Proteins play a central role in biological processes, and understanding their conformational variability is crucial for unraveling their functional mechanisms. Recent advancements in high-throughput technologies have enhanced our knowledge of protein structures, yet predicting their multiple conformational states and motions remains challenging. This study introduces Dimensionality Analysis for protein Conformational Exploration (DANCE) for a systematic and comprehensive description of protein families conformational variability. DANCE accommodates both experimental and predicted structures. It is suitable for analysing anything from single proteins to superfamilies. Employing it, we clustered all experimentally resolved protein structures available in the Protein Data Bank into conformational collections and characterized them as sets of linear motions.

The resource facilitates access and exploitation of the multiple states adopted by a protein and its homologs. Beyond descriptive analysis, we assessed classical dimensionality reduction techniques for sampling unseen states on a representative benchmark. This work improves our understanding of how proteins deform to perform their functions and opens ways to a standardised evaluation of methods designed to sample and generate protein conformations.

### 12 Introduction

Proteins orchestrate all biological processes, and their malfunctions often result in disease. In recent years, high-throughput 13 technologies have greatly improved our knowledge of their amino acid sequences and 3D shapes<sup>1-4</sup>. While reaching the 14 single-structure frontier<sup>5</sup>, these advances have also highlighted the complexities of how proteins move and deform to carry 15 out their biological functions<sup>6,7</sup>. They have stimulated a renewed interest in the modeling of protein and protein complex 16 multiple conformational states<sup>8</sup>. In particular, the success of the protein structure prediction neural network AlphaFold<sup>29</sup> 17 has inspired innovative strategies for modifying or repurposing it toward exploring protein conformational space. These 18 approaches involve forced sampling<sup>10</sup>, modulation of input multiple sequence alignment content and depth<sup>11,12</sup>, or guidance 19 with state-annotated templates<sup>13,14</sup>. Although they have achieved promising results for specific protein families, systematic 20 assessments have revealed limitations<sup>15,16</sup>. In addition, studies sampling from low-dimensional representations or manifolds 21 learned from observed or simulated conformations<sup>17-19</sup> have underscored the difficulty in predicting new, completely unseen 22 states and the importance of high-quality data for training or benchmarking. 23 Experimental techniques like X-ray crystallography, cryogenic-electron microscopy (cryo-EM), and nuclear magnetic

24 resonance spectroscopy (NMR) are essential for capturing protein functional states<sup>6,20</sup>. The Protein Data Bank (PDB)<sup>4</sup> offers 25 access to multiple structural states for various proteins, solved independently in different conditions, oligomeric states, and with 26 diverse cofactors and molecular partners. Researchers have actively engaged in efforts to collect, cluster, curate, represent, 27 visualise, and functionally annotate these states 20-23. These endeavours have provided valuable insights into the biologically 28 meaningful conformational space for specific protein families such as protein kinases<sup>24</sup>, RAS isoforms<sup>25</sup>, ABC (ATP Binding 29 Cassette) transporters<sup>26</sup>, and G-protein coupled receptors  $(GPCRs)^{27}$ . However, producing or validating functional annotations 30 for structural states involves a substantial amount of manual intervention. Despite the wealth of experimentally resolved protein 31 conformational variability, its full exploitation remains an ongoing challenge. 32

Ideally, one would like to comprehensively describe protein conformational variability with low-dimensional representations or manifolds amenable to visualisation and interpretation. Principal Component Analysis (PCA) serves as a convenient and robust means to reduce the dimensionality of a dataset, capturing maximum variability<sup>28,29</sup>. The principal components extracted from a conformational ensemble define 3D directions for every atom, and motions along them allow navigating the conformational space<sup>30</sup>. PCA has proven useful for extracting structural transitions from sparse disconnected low-energy structural states<sup>31-36</sup>. Unlike more complex non-linear dimensionality reduction techniques, it offers the advantage of not
 depending on numerous adjustable parameters and provides a straightforward geometrical interpretation.

<sup>40</sup> Here, we describe a PDB-wide analysis of protein conformational variability across various levels of sequence homology.

- <sup>41</sup> Our fully-automated computational pipeline, named Dimensionality Analysis for protein Conformational Exploration (DANCE),
- <sup>42</sup> systematically compiles collections of aligned protein conformations and extracts their principal components. We interpret
- the representation space defined by the main principal components as the *linear motion manifold* underlying the observed
- 44 conformations. We provide estimates of the intrinsic dimensionality of these motion manifolds. To assess generative methods,
- <sup>45</sup> we introduce a benchmark set comprising ten conformational collections representing therapeutic targets with substantial
- <sup>46</sup> functional transitions. Additionally, we provide baseline performances from classical linear and non-linear manifold learning
   <sup>47</sup> techniques.
- <sup>48</sup> DANCE is versatile, handling both experimental and predicted structures with varying amino acid sequences. It adopts

<sup>49</sup> an unbiased approach, avoiding predetermined protein or domain definitions when building the conformational collections.

- 50 Considering the complete context of input protein chains enables a thorough examination of inter-domain motions. Furthermore,
- <sup>51</sup> DANCE accommodates uncertainty from unresolved protein regions without assuming potential conformations. It introduces a
- <sup>52</sup> weighting scheme to mitigate the imbalanced coverage of variables.

We provide several databases of conformational collections representing the whole PDB as well as detailed information about the benchmark on Figshare. In addition, DANCE's source code is available at: https://github.com/ 54 PhyloSofS-Team/DANCE.

### 56 Methods

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### 57 Overview of DANCE

<sup>58</sup> DANCE takes as input a set of protein 3D structures (in Crystallographic Information File or CIF format) and outputs a <sup>59</sup> set of protein- or protein family-specific conformational collections or ensembles (in CIF of PDB format). It first clusters

and superimposes the input structures based on the similarities found in their corresponding amino acid sequences. It then

determines the set of principal components sufficient to explain the variability observed within each conformational ensemble.

<sup>62</sup> The algorithm unfolds in six main steps depicted in **Fig. 1**.

a- Extraction of sequences. The first step extracts the one-letter amino acid sequences of all polypeptidic chains contained in the input CIF files. In case of multiple models, DANCE retains only the first one. The names of the residues with resolved 3D coordinates are taken from the \_atom\_site.label\_comp\_id column. Residues missing from the protein structure are included as lowercase letters in the sequence if they are defined in the \_entity\_poly\_seq category. This information will help in clustering and aligning the sequences (see below). Otherwise, they are replaced by the "X" symbol. The "X" symbol is also used for unknown amino acid types and for modified amino acids without a close natural neighbour. Sequences comprising less than 5 non-"X" residues are then filtered out.

- **b- Clustering of the sequences.** DANCE clusters sequences using MMseqs2<sup>37</sup>. The users can choose the desired levels of sequence similarity and coverage, both set to 80% by default. The coverage is bidirectional by default. This step outputs a TSV file specifying the clusters.
- c- Multiple sequence alignments. DANCE then aligns the sequences within each cluster using MAFFT<sup>38</sup> with default
   parameters. It further removes all the columns containing only Xs or gaps, and reorders the sequences according to their
   PDB codes.
- **d Extraction of structures.** DANCE extracts 3D coordinates of the backbone atoms N, C, C $\alpha$ , and the O atom, of all polypeptidic chains contained in the input CIF files. It reconstructs missing O atoms based on the other atom's coordinates. It disregards residues with missing backbone atoms and chains shorter than 5 residues.
- e- Generation of the conformational collections. DANCE then uses the sequence clusters defined in (b) to group 79 conformations and the residue matching provided by (c) to superimpose them. The superimposition puts their centers of 80 mass to zero and then aims at determining the optimal least-squares rotation matrix minimizing the Root Mean Square 81 Deviation (RMSD) between any conformation and a reference conformation (see below). This is achieved through the 82 ultrafast Quaternion Characteristic Polynomial method<sup>39,40</sup>. The users can choose to account for all the atoms in the 83 superimposition, or only the C $\alpha$  atoms. Optionally, the users can filter out the conformations with too few (less than 5 by 84 default) residues aligning to the reference. As a post-processing step, DANCE reduces structural redundancy. Namely, 85 it removes any conformation A deviating by less than  $rms_{cut}$  Å from another one B, provided that the sequence of A is 86 identical to or included in that of B. The value of rms<sub>cut</sub> is 0.1 Å by default and is customizable by the users. Finally, 87

DANCE saves the conformational ensemble as a multi-model file in PDB or CIF format. Notice that the models can
 display different amino acid sequences. DANCE also outputs the corresponding multiple sequence alignments (MSA) in

<sup>90</sup> FASTA format, and the matrix of all-to-all pairwise RMSDs.

• **f**- **Extraction of linear motions.** DANCE performs PCA on the 3D coordinates from each collection. This dimensionality reduction technique identifies orthogonal linear combinations of the variables, namely the Cartesian coordinates, maximally explaining their variance (see below). These linear combinations, which we refer to as principal components or PCA modes, represent directions in the 3D space for every atom. Deforming the protein structure using these components produce motions that connect the conformations observed in the collection. For the sake of simplicity, we directly refer to the principal components as to *linear motions*, although they may not represent actual physical motions undergone by the protein. Furthermore, we estimate the *intrinsic dimensionality* of the linear motion manifold underlying an ensemble's conformational variability as the number of principal component explaining essentially all its positional variance. The higher the dimensionality – the more complex the linear motions.

### 100 Choosing a reference

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We choose the reference conformation for the superimposition as the one with the amino acid sequence most representative of the MSA. For this, we first determine the consensus sequence  $s^*$  by identifying the most frequent symbol at each position. We consider "X" symbols as equivalent to gaps. Hence, each position is described by a 21-dimensional vector giving the frequencies of occurrence of the 20 amino acid types and of the gaps. In case of ambiguity, we prefer an amino acid over a gap and a more frequent amino acid over a less frequent one. Then, we compute a score for each sequence *s* in the MSA reflecting its similarity to  $s^*$  and expressed as,

$$\operatorname{score}(s) = \sum_{i=1}^{P} \sigma(s_i, s_i^*), \tag{1}$$

where *P* is the number of positions in the MSA and  $\sigma(s_i, s_i^*)$  is the substitution score between the amino acid  $s_i$  at position *i* in sequence *s* and the consensus symbol  $s_i^*$  at position *i*. We use the substitution matrix BLOSUM62 and we set the gap score to  $\min_{a,b}(\sigma(a,b)) - 1 = -5$ . MAFFT also uses BLOSUM62 for generating the MSAs.

### Judging the quality of the MSA

We compute the identity level of an MSA as the average percentage of sequence pairs sharing the same amino acid in a column, and the coverage as the percentage of positions having less than 20% of gaps. In addition, we evaluate the global quality of the MSA with a sum-of-pairs score, with  $\sigma_{match} = 1$  and  $\sigma_{mismatch} = \sigma_{gap} = -0.5$ . We normalise the raw sum-of-pairs scores by dividing them by the maximum expected values. The final score for an MSA is thus expressed as,

$$score_{rel}(MSA) = \frac{score(MSA)}{\binom{n}{2}L_{eff}},$$
(2)

where is the raw MSA score, n is the number of chains or sequences, and  $L_{eff}$  is the effective length of the MSA, computed as,

$$L_{eff} = \max_{s \in \mathscr{S}} \sum_{i=1}^{L(s)} \mathbb{I}\{s_i \in \mathscr{A}\},\tag{3}$$

where  $\mathscr{S}$  is the set of sequences comprised in the MSA, L(s) is the length of the aligned sequence *s*, and  $\mathscr{A}$  is the 20-letter amino acid alphabet (*e.g.*, excluding gap characters).

### 113 Extracting linear motions

The Cartesian coordinates of each conformational ensemble can be stored in a matrix *R* of dimension  $3m \times n$ , where *m* is the number of positions in the associated MSA and *n* is the number of conformations. Each position is represented by a C- $\alpha$  atom. We compute the covariance matrix as,

$$C = \frac{1}{n-1} R^c (R^c)^T = \frac{1}{n-1} (R - \bar{R}) (R - \bar{R})^T,$$
(4)

where  $\bar{R}$  is obtained by averaging the coordinates over the conformations. Alternatively, the users can choose to center the data on the reference conformation. The covariance matrix is a  $3m \times 3m$  square matrix, symmetric and real.

The PCA consists in decomposing C as  $C = VDV^T$  where V is a  $3m \times 3m$  matrix where each column defines an eigenvector or a PCA mode that we interpret as a linear motion. D is a diagonal matrix containing the eigenvalues. The sum of the eigenvalues  $\sum_{k=1}^{3m} \lambda_k$  amounts to the total positional variance of the ensemble. The portion of the total variance explained by the the eigenvector or linear motion is estimated as  $\frac{\lambda_k}{\sum_{k=1}^{3m} \lambda_k}$ .

In addition, we estimate the collectivity  $^{41,42}$  of the *k*th eigenvector as,

$$\operatorname{coll}(\mathbf{v}_k) = \frac{1}{m} \exp\left(-\sum_{i=1}^{3m} v_{ki}^2 \log v_{ki}^2\right).$$
(5)

If  $coll(\mathbf{v}_k) = 1$ , then the corresponding motion is maximally collective and has all the atomic displacements identical. In case of an extremely localised motion, where only one single atom is affected, the collectivity is minimal and equals to 1/m.

We also apply PCA to the correlation matrix computed by normalising the covariance matrix as,

$$\operatorname{Cor}_{i,j} = \frac{C_{i,j}}{\sqrt{C_{i,i}}\sqrt{C_{j,j}}}.$$
(6)

In that case, the sum of the eigenvalues  $\sum_{k=1}^{3m} \lambda_k$  amounts to 1.

### 123 Handling missing data

As stated above, the conformations in a collection may have different lengths reflected by the introduction of gaps in the associated MSA. We fill these gaps with the coordinates of the conformation used to center the data (average conformation, by default). In doing so, we avoid introducing biases through reconstruction of the missing coordinates. Moreover, this operation results in low variance for highly gapped positions, thus limiting their contribution to the extracted motions. To go further and explicitly account for data uncertainty, we implemented a weighting scheme. Specifically, DANCE assigns confidence scores to the residues and include them in the structural alignment step and the PCA. The confidence score of a position *i* reflects its coverage in the MSA,  $w_i = \frac{1}{n} \sum_{S} \mathbb{1}_{a_i^S \neq "X"}$ , where "X" is the symbol used for gaps. The structural alignment of the *j*th conformation onto the reference conformation amounts to determining the optimal rotation that minimises the following function<sup>43</sup>,

$$E = \frac{1}{\sum_{i} w_{i}} \sum_{i} w_{i} (r_{ij}^{c} - r_{i0}^{c})^{2},$$
(7)

where  $r_{ij}^c$  is the *i*th centred coordinate of the *j*th conformation and  $r_{i0}^c$  is the *i*th centred coordinate of the reference conformation. The resulting aligned coordinates are then multiplied by the confidence scores prior to the PCA.

### 126 Implementation details

We implemented DANCE in C/C++ and Python. It relies on the C++ GEMMI library<sup>44</sup> to parse the CIF files and manipulate the structures. It runs MMseqs2 through the following command: *cluster DB clusterDB tmp –cov-mode 0 -c* \$*cov –min-seq-id* \$*id*. It launches MAFFT with the options *auto*, *amino* and *preservecase*. The multiple sequence alignment and structure superimposition steps are parallelized. For the PCA, we use the singular value decomposition (SVD) implemented in NumPy<sup>45</sup> on the *R* matrix directly. SVD is computationally more advantageous when  $3m \gg n$ , which is typically the case of our data,

since we only compute the required number of n components.

### 133 Application and extension of DANCE

DANCE is applicable to experimental 3D structures as well as predicted 3D models, as long as they comply with the CIF standards.

### 136 Describing conformational variability over the whole PDB

We applied DANCE to all 748 297 protein chains with experimentally resolved 3D structures available in the PDB, as of June 137 2023. We downloaded all the PDB entries in CIF format from the  $RCSB^{46}$ . We replaced the raw CIF files with their updated 138 and optimised versions from PDB-REDO whenever possible<sup>47</sup>. It took about 2.25 hours to run DANCE on the whole PDB 139 on a desktop computer with Intel Xeon W-2245 @ 3.90GHz and 32Go of RAM (Table S1). The most time consuming steps 140 are the extraction and superimposition of the 3D structures to create the conformational ensembles. We ran DANCE at eight 141 different levels of sequence similarity, designated as  $l_{cov}^{id}$ , where *id* and *cov* are the sequence identity and coverage thresholds, 142 correspondingly, and range from 50 to 80%. For investigating how the ensembles transformed across levels, we focused on the 143 18 616 conformational ensembles detected in the most relaxed set up, namely at 30% identity and 50% coverage  $(l_{30}^{50})$ . For each 144 ensemble, we extracted its reference protein chain and we traced back the conformational ensembles to which it belonged upon 145 progressively applying stricter thresholds. 146

### 147 Focusing on the ABC superfamily

We extended DANCE usage beyond the single-chain and sequence-similarity paradigms to describe the conformational 148 variability of ABC (ATP Binding Cassette) transporters. We retrieved a set of 354 ABC protein experimental 3D structures 149 from https://abc3d.hegelab.org<sup>26</sup>. They correspond to functionally relevant states annotated as biological units in 150 the PDB. In most of these structures, several polypeptidic chains, typically 2 or 4, encode the two nucleotide-binding domains 151 (NBDs) and two transmembrane domains (TMDs) of the ABC architecture. In addition, some structures contain several ABC 152 protein copies or some ABC protein cellular partners (small molecules, substrate peptides, interacting proteins). We chose 153 the murine ABC transporter P-glycoprotein (5KOYA) as reference for the subsequent analysis. Its 1182-residue long single 154 polypeptidic chain the full-length transporter architecture. 155 To cope with the high sequence divergence of the ABC superfamily, we relied on structural similarity for grouping and 156

matching the ABC conformations. Specifically, we used the method Foldseek<sup>48</sup> to identity structures sharing significant 157 similarity with the reference and align them. We performed a first screen by querying the reference against all individual chains 158 (1 244 in total) and defined significant hits as those with an e-value lower than 10.0. Then, for each structure, we estimated 159 an upper bound on its coverage of the reference by summing up the reference residue ranges appearing in the alignments 160 associated with its significant hits. We filtered out the structures with coverage upper bounds lower than 90%. We performed 161 a second screen by querying the reference against the 209 remaining structures defined as monomers by concatenating their 162 chains. We identified two structures (5NIK, 5NIL) spanning less than 90% of the reference. Permuting their chains did not 163 increase their coverage and thus we removed them. To further detect potentially suboptimal chain orderings, we computed 164 reference to target residue span ratios. We identified one structure, namely 7AHD, with a highly imbalanced ratio of 1.6. Such 165 a high value is indicative of large parts of the reference that could not be aligned to the target structure. Permuting the four 166 chains (A,B,C,D) of 7AHD into (A,D,B,C) led to a more balanced ratio of 0.86. We did not observe discrepancies for other 167 structures and thus we retained their original chain ordering. Finally, we removed the structures with low-quality alignments, 168 *i.e.*, with more than 200 gaps or with a continuous gapped region of more than 60 positions. 169

Among the 195 structures finally selected, 4F4C, 7SHN and 7AHD contained unknown or unrecognized amino acids which 170 we removed. We ran Foldseek one more time to generate a structure similarity-based multiple sequence alignment centred on 171 the reference 5KOYA. We trimmed the alignment and the 3D structures by removing the residues inserted with respect to the 172 reference. We gave the trimmed alignment and 3D coordinate files as input to DANCE, starting directly from step d (see the 173 overview of DANCE algorithm above). For consistency and comparison purposes, we asked DANCE to center the data on 174 the reference. To mitigate the impact of potential alignment errors, we applied weights reflecting position-specific confidence 175 scores (see above, *Handling missing data*). DANCE structural redundancy reduction step removed 7 conformations, resulting 176 in an ensemble of 188 conformations. 177

We compared this ensemble with those generated by DANCE default sequence similarity-based end-to-end procedure applied to the whole PDB. More specifically, we took the ensembles generated at  $l_{80}^{80}$  and  $l_{50}^{30}$  and containing 5KOYA and we rebuilt them with DANCE, applying the 5KOYA centering and the uncertainty weighting scheme. We estimated the similarity between the ensembles' motion subspaces as the Root Mean Square Inner Product (RMSIP)<sup>49,50</sup>. The latter measures the overlap between all pairs of the *l* first PCA modes and is defined as,

$$\text{RMSIP} = \sqrt{\frac{1}{l} \sum_{i=1}^{l} \sum_{j=1}^{l} (\mathbf{v}_{i}^{\mathscr{S}_{\mathscr{A}}} \cdot \mathbf{v}_{j}^{\mathscr{S}_{\mathscr{A}}})^{2}},$$
(8)

where  $\mathbf{v}_i^{\mathscr{S}_{\mathscr{A}}}$  and  $\mathbf{v}_j^{\mathscr{S}_{\mathscr{A}}}$  are the *i*th and *j*th PCA modes extracted from the conformational ensembles  $\mathscr{S}_{\mathscr{A}}$  and  $\mathscr{S}_{\mathscr{B}}$ , and *l* is the number of modes considered for the comparison. Moreover, we monitored the distance between the geometric centres of the two NBDs defined by the C- $\alpha$  atoms of residues numbered 346-596 and 929-1182, respectively, in the reference 5KOYA.

### 181 Benchmarking for the generation of unseen conformations

### 182 Linear PCA

We further investigated whether the extracted principal components could be useful to predict unseen conformations. Given a set of *l* PCA modes computed from the coordinates *R*, we generate a new conformation  $\mathbf{r}_{nred}^*$  as,

$$\mathbf{r}_{\mathbf{pred}}^* = \mathbf{p}^* V_l^T + \bar{\mathbf{r}},\tag{9}$$

where the matrix  $V_k \in \mathbb{R}^{3m \times l}$  contains the modes,  $\mathbf{\bar{r}} \in \mathbb{R}^{3m}$  is the average conformation, and  $\mathbf{p}^* \in \mathbb{R}^l$  is a point in the *l*-dimensional representation space defined by the modes. The coordinates of  $\mathbf{p}^*$  specify the amplitudes of the modes.

#### Nonlinear kernel PCA 185

The manifold underlying our data is a priori non-linear. This motivated us to investigate whether non-linear methods could 186

achieve better reconstructions than linear PCA. We focused on the widely used kernel Principal Component Analysis (kPCA)<sup>51</sup>. 187

The intuition behind kPCA is to map the input data points to a higher dimensional space where they will be linearly separable by 188

a classical PCA. The mapping function  $\phi : \mathbb{R}^{3m} \to \mathbb{R}^M$  is not known. Instead of explicitly calculating it, we use a kernel function 189  $k(\mathbf{r}_i,\mathbf{r}_i) = \phi(\mathbf{r}_i)^T \phi(\mathbf{r}_i)$ , where  $\mathbf{r}_i$  and  $\mathbf{r}_i$  are two conformations. We chose the radial basis function (RBF) kernel, defined as

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 $k(\mathbf{r}_i, \mathbf{r}_j) = e^{-\frac{d(\mathbf{r}_i, \mathbf{r}_j)^2}{2\sigma^2}}$ , where  $d(\mathbf{r}_i, \mathbf{r}_j)$  is the Euclidean distance between the two conformations  $\mathbf{r}_i$  and  $\mathbf{r}_j$ . We explored different values of the hyperparameter  $\sigma$ . For sufficiently small values, *i.e.*,  $\frac{1}{2\sigma^2}d(\mathbf{r}_i, \mathbf{r}_j)^2 \ll 1$ , the RBF kernel becomes effectively linear, 191 192 since, in this case,  $k(\mathbf{r}_i, \mathbf{r}_j) \approx 1 - \frac{1}{2\sigma^2} d(\mathbf{r}_i, \mathbf{r}_j)^2$ . 193

Thus, given the input coordinates R representing n conformations, we computed the corresponding RBF kernel matrix K of dimension  $n \times n$  and decomposed it using the classical PCA. The resulting principal components  $\{v_1, v_2, ..., v_n\}$  can then be expressed as

$$\mathbf{v}_j = \sum_{i=1}^n a_{ji} \phi(\mathbf{r}_i). \tag{10}$$

They allow extracting nonlinear features but they cannot be combined straightforwardly to generate new conformations. Instead, 194 for generative purposes, we need to learn an inverse transform function that maps points in the *l*-dimensional representation 195 space defined by the components back to the input space. This problem is known as the pre-image problem. To solve it, we used 196 kernel ridge regression of the input coordinates R on their low-dimensional projections in the representation space as described 197 in<sup>52,53</sup> and implemented in the scikit-learn Python library<sup>54</sup>. The contribution of the L2-norm regularisation is controlled 198 through the hyperparameter  $\alpha$ . More technically,  $\alpha$  connects the squared L2-norm between a point in the representation space 199 and its reconstruction with the squared L2-norm of the kernel weights used for the reconstruction. 200

#### Leave-one-cluster-out cross-validation procedure 201

We assessed the predictive performance of PCA and kPCA with a *leave-one-out* cross-validation procedure. Since the 202 conformations are not evenly distributed within an ensemble, we grouped them into clusters prior to the evaluation. We 203 performed the clustering in the l-dimensional PCA representation space, where l is the minimal number of linear components 204 sufficient to explain 90% of the ensemble's total positional variance. We used the k-means clustering<sup>55</sup> with k = l + 2. 205

Given a clustered ensemble, we systematically tested the ability of the principal modes inferred from l+1 clusters to predict the conformations belonging to the held-out cluster. We reconstructed each test conformation  $\mathbf{r}^*$  from its projection  $\mathbf{p}^*$  in the *l*-dimensional representation space. For the classical PCA, we computed the projection as,

$$\mathbf{p}^* = (\mathbf{r}^* - \bar{\mathbf{r}}) V_l. \tag{11}$$

For the kPCA, the projection onto the principal component  $v_i$  is expressed as,

$$\phi(\mathbf{r}^{*})\mathbf{v}_{\mathbf{j}} = \sum_{i=1}^{n} a_{ji}\phi(R)^{T}\phi(\mathbf{r}^{*}) = \sum_{i=1}^{n} a_{ji}K(R,\mathbf{r}^{*}).$$
(12)

We evaluated the reconstruction error as the RMSD between the predicted conformation  $\mathbf{r}_{\text{pred}}^*$  and the original conformation  $\mathbf{r}^*$ . 206

#### Distance to the training set 207

We estimated the difficulty of reconstructing a given conformation by computing its distance to the convex hull defined by the 208 conformations used for training in the *l*-dimensional representation space. Setting the number of clusters in the training set to 209 l+1 ensures that the convex hull will be a polytope of dimension at least l. For instance, in 1 dimension, we need at least 2 210 affine-independent points to define a 1-polytope. The explicit computation of the convex hull of n points in l dimensions is 211 an operation whose complexity is of the order of  $O(n^{l/2})^{56}$  and rapidly becomes computationally infeasible as the value of 212 *l* increases. Nevertheless, the calculation of the distance of a given point to the hull does not require computing the convex 213 hull explicitly and is a much simpler computational problem. It can be solved in quasilinear time with quadratic programming 214 (QP). Here, we used the efficient and exact QP simplex solver proposed in<sup>57</sup> and implemented in the Computational Geometry 215 Algorithms Library (CGAL)<sup>58</sup>. It takes advantage of the low dimensionality of the representation space by observing that the 216 closest features of two *l*-polytopes are always determined by at most l + 2 points. 217

In order to compare distances across systems of different sizes, we scale them by the number of positions m,

$$d^{norm} = \frac{d}{\sqrt{m}}.$$
(13)



**Figure 1. Outline of the study.** Our approach, DANCE, exploits both amino acid sequences and 3D coordinates. We applied it to all experimentally determined protein-containing 3D structures from the PDB. Alternatively, users can provide a custom set of experimental structures or predicted models. DANCE first concentrates on sequences. It extracts them from the input structures (A) and clusters them with MMseqs2 based on user-defined similarity and coverage thresholds (B). For each cluster, It generates a multiple sequence alignment using MAFFT (C). It then extracts all 3D coordinates (D), groups the conformations according to the clusters identified in B and superimposes them to generate conformational ensembles (E). The superimposition aims at minimizing the Root Mean Square Deviation to a chosen reference, using the alignments produced by C for mapping the residues. The examples of the bacterial enzymes adenylate kinase (in grey, reference PDB code: 1AKEA) and MurD (in blue, 1E0DA), and the murine ABC transporter P-glycoprotein (5KOYB) are depicted. The arrows indicate adenylate kinase's main motion. The horizontal lines behind the P-glycoprotein indicate the boundaries for the membrane bilayer. Finally, DANCE summarises conformational diversity through Principal Component Analysis (F). We further assessed the ability of classical manifold learning techniques to reconstruct and extrapolate conformations.

This normalisation also allows relating distances in the representation space with RMS deviations in the 3D Cartesian space.
 Indeed, let us consider an ensemble of conformations exhibiting a purely one-dimensional motion. Any two conformations
 distant by an RMSD of 1 Å in the original 3D space will be separated by a normalised distance of 1 Å in the one-dimensional
 representation space.

### 222 Results

<sup>223</sup> We used DANCE to chart the experimentally resolved conformational diversity of protein families (**Fig. 1**). We explored eight

- levels of sequence similarity (*sin*) and coverage (*cov*), denoted as  $l_{cov}^{sim}$ , to group the ~750K chains included in the PDB as of
- June 2023 (Fig. S1A and Table S2). In the most conservative set up, namely  $1_{80}^{80}$ , less than 3% of the conformations remain

isolated (Fig. S1A, singletons). Most of the conformational collections (or ensembles) are associated with multiple sequence 226

alignments of high quality across all levels (Fig. S1B). Sequence identity and coverage are more widely distributed in more 227

relaxed conditions, but the median values always remain very high, above 0.95 (Fig. S1C-D). 228

#### Experimentally resolved conformations lie on low-dimensional manifolds 229

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Only one or two linear principal components suffice to explain almost half of the ensembles' conformational diversity (Fig. 2A). We interpret these components as directions of motion, and by simplification, we will denote them as linear motions in the following (see *Methods*). In the overwhelming majority of cases, less than eight linear motions explain more than 90% of the total positional variance. These observations hold true across all sequence identity and coverage levels. They indicate that the 233 conformational states captured by experimental techniques for a protein or a protein family lie on a low-dimensional manifold. This low dimensionality is only partially determined by the cardinality of the ensembles (Fig. S2A-B). Almost 30% of the most 235 highly populated ensembles (>50 conformations) detected at  $l_{80}^{80}$  can be comprehensively described with less than three linear 236 motions (Fig. S2C). This proportion increases up to 46% in the most relaxed conditions, namely at  $l_{50}^{30}$  (Fig. S2D).

The bacterial adenylate kinase gives an example of a one-dimensional motion underlying its 42 conformations (Fig. 1E. 238 in grey). One can easily classify the conformations by visual inspection into two main states, open and closed, deviating by 239 about 7 Å. The bacterial enzyme MurD (Fig. 1E, in blue) and the murine ABC transporter P-glycoprotein (Fig. 1E, in orange) 240 also exhibit low-dimensional opening-closing motions. In particular, the P-glycoprotein's collection reveals a rich spectrum of 24 intermediate conformations between the open and closed forms (Fig. 1E, in orange). The main motion involves about 70% 242 of the protein and modulates the volume of the transporter's internal cavity within the lipid bilayer up to over  $6,000 \text{ Å}^{359}$ . It 243 explains about 80% of the total positional variance on its own. The remaining variability is mostly due to rotations of the 244 nucleotide binding domains with respect to the transmembrane helical bundles and to loop deformations. 245

#### A few protein families display huge conformational expansion upon relaxing the sequence selection crite-24F ria 247

To investigate how the conformational ensembles transformed with sequence similarity, we systematically backtracked the 248 18 616 representative protein chains identified at  $l_{50}^{30}$  across more stringent levels (see *Methods*). The fragment antigen-binding 249 regions display the largest growth between the most stringent and most relaxed sequence selection criteria (Fig. 2). For 250 instance, while the Fab6785 light chain's ensemble at  $I_{80}^{80}$  comprises a bit less than 300 conformations, it expands up to over 251 12 500 conformations at  $l_{50}^{30}$  (Fig. 2B, PDB id: 4QHUH). With the largest number of conformations at  $l_{80}^{80}$ , the HIV-1 capsid 252 protein's ensemble however displays a relatively limited expansion across the different levels, from 3 334 to 3 391 (Fig. 2B, 253 3J345). Bovine trypsin and its close homologs give an example of an extensively characterized subfamily, with 470 different 254 conformations detected at  $l_{80}^{80}$ . This ensemble expands by more than 5 folds, aggregating different serine proteases, upon relaxing the criteria to  $l_{50}^{30}$  (**Fig. 2B**, PDB id: 1TAWA). Likewise, the Beta-2-microglobulin and its close homologs have a large body of 1 465 conformations at  $l_{80}^{80}$ , growing further up to 2 025 conformations at  $l_{50}^{30}$  by including other immunoglobulins (**Fig. 2B**, 7MX4B). By contrast, the reconstructed ancestral tyrosine kinase AS, a common ancestor of Src and Abl, has only 2 255 256 257 258 conformations available in the PDB and no close homologs. At  $l_{50}^{30}$ , it serves as representative for a huge ensemble of over 259 4 000 protein kinase conformations (Fig. 2B, 4UEUA). Apart from these over-represented protein families or superfamilies, the 260 ensembles generally gain only a few conformations, with a median value of 4. 26

#### Family expansion may lead to an apparent motion simplification 262

As an ensemble grows, the gained conformations may lie on the same motion manifold, defined by the subset of principal 263 components explaining the variance, or give rise to new motions represented by new components (Fig. 2C). The bacterial 264 long-chain flavodoxin exemplifies the second scenario (Fig. 2D-F, in black). At  $l_{80}^{80}$ , it undergoes a one-dimensional motion 265 describing the transition between a compact state and a partially unfolded conformation (Fig. S3). Upon relaxing sequence 266 similarity to  $l_{50}^{30}$ , the ensemble roughly doubles in size (Fig. 2F) and the newly added conformations exhibit complex 267 deformations of the FMN binding pocket. As a result, five more linear motions are required to explain the positional variance 268 (Fig. 2D). Hence, in this case, the motions get more complex when considering more distant homologs. 269

The emergence of new motions does not however systematically lead to an increased motion complexity. The murine MCL1 270 gives an illustrative example of apparent motion simplification upon expansion (Fig. 2D-F, in red, and Fig. 2G). At  $l_{80}^{80}$ , almost 271 30 components are needed to explain the variability observed over the couple of hundreds conformations in the ensemble. They 272 represent local deformations of the inter-helical loops and the extremities (Fig. 2G and Fig. S3). Extending the ensemble to 273 distant members of the Bcl-2 family brings in about 50 new conformations (Fig. 2F). They reveal a new extended state the 274 protein BAX adopts upon assembling into domain-swapped dimers<sup>60</sup>. The large amplitude transition between the compact 275 conformation and the extended one takes a big part in the variance, resulting in a drastically reduced motion complexity (Fig. 276 2D). The benzaldehyde lyase BAL gives another example (Fig. 2D-F, in blue) where the transition to a new state, adopted 277



**Figure 2.** Evolution of protein conformational diversity across sequence similarity levels. A. Proportion of conformational ensembles requiring *n* linear PCA modes to explain 90% of their total positional variance, with *n* varying from 1 to 8. The number of modes *n* is an indicator of motion complexity. Singletons and pairs are excluded. **B.** Cumulative distribution of the number of conformations gained from the most stringent level, namely  $1_{80}^{80}$ , with 80% sequence similarity and coverage, to the most permissive one,  $1_{50}^{30}$ , with 30% similarity and 50% coverage. The 3D structures of the reference protein chains are depicted for a few ensembles. **C.** Comparison of motion complexity between the most stringent and most relaxed set ups. We considered only the cases where the ensemble at  $1_{50}^{30}$  is bigger than the corresponding one at  $1_{80}^{80}$ . Singletons and pairs are excluded. **D-G.** Detailed evolution of three ensembles marked by colored dots in panel C. **D.** Motion complexity expressed as a number of modes. The names and PDB codes of the reference chains are indicated. **E.** Motion amplitude, measured as the maximum RMSD between any two conformations (in Å). **F.** Conformational collection size. **G.** Conformational diversity observed for the Bcl-2 family. On the top left, the 54 conformations comprised in the MCL1 ensemble at  $1_{80}^{80}$ . At the bottom right, the 218 additional conformations at  $1_{50}^{30}$ . The color code indicates the position in the sequence, from the N-terminus in blue to the C-terminus in red.

by the distant homolog actinobacterial 2-hydroxyacyl-CoA lyase<sup>61</sup>, dominates the variance (**Fig. S3**). The conformational variability transforms from small (<1Å) seemingly random fluctuations to a one-dimensional motion.

Overall, about a third of the ensembles undergo an apparent motion simplification upon expansion (**Fig. 2C** and **Fig. S4A**). They likely represent protein families where distant homologs exhibit novel distinct states. The larger the deviations of these novel states with respect to the other ones, the higher the contribution of the corresponding motions to the variance. To mitigate this variance-dependent effect, we repeated the analysis on the correlation matrix. The latter estimates the extent to which the

residues move in the same direction, regardless of the magnitude of their displacements. We found that the motion complexity relification does not merch accelerate the same direction of the same di

still decreases in over 20% of the ensembles (**Fig. S4B**). This result indicates that motion simplification does not merely reflect larger transitions "hiding" smaller rearrangements. A substantial fraction of protein families show evidence of more concerted

residue movements between more distant homologs.

### 288 Beyond single chains and sequence similarity, the ABC superfamily as a case study

We explored the possibility of using DANCE to chart the conformational variability of remote homologs with low sequence similarity and variable chain composition. We focused on the ABC (ATP Binding Cassette) transporter superfamily. The ABC architecture comprises two nucleotide-binding domains (NBDs) and two transmembrane domains (TMDs) encoded by one or several polypeptidic chains (**Fig. 3A**). The NBDs are highly conserved across species and families, whereas the TMDs exhibit various scaffolds associated with heterogeneous transport functions<sup>26</sup>. We considered a collection of a few hundreds ABC protein experimental 3D structures<sup>26</sup>, taking the single-chain murine P-glycoprotein as reference (**Fig. 3A**, 5KOYA).

We bypassed DANCE sequence extraction, clustering and alignment steps and directly gave it a pre-computed alignment 295 built from structural similarities as input (see Methods). Relying on structure rather than sequence similarity and considering 296 various oligomeric states provided a more comprehensive description of ABC transporters' functional motions and states (Fig. 297 3 and Movies S1-2). The resulting ensemble comprises 188 conformations encompassing 295 protein chains, some of which 298 have sequence identity below 30% or coverage lower than 50% (Fig. 3A). A set of 25 linear motions are required to explain 299 the positional variance. By comparison, the sequence similarity-based 5KOYA-containing collection generated by DANCE at 300  $1_{50}^{30}$  contains only 71 conformations explained by only four linear motions. These motions are essentially identical to those 301 extracted from the 61 conformations at  $l_{80}^{80}$  (Fig. 3B, RMSIP = 0.99). 302

Despite having different motion complexities, the sequence- and structure-based conformational collections have largely 303 overlapping motion subspaces (Fig. 3B, RMSIP  $\sim 0.7$ ). In particular, they all share the same most contributing motion 304 describing the transition between the transporter inward-closed and inward-open forms (Fig. S5). This functional transition 305 controls the substrate access to the transporter's central binding pocket. It explains 45 to 70% of the variance on its own and 306 involves over two-thirds of the residues. The structure similarity-based collection represents a quasi-continuum of increasingly 307 open states (Fig. 3C, in blue, and Movie S1) between two extreme dimeric forms, one from the human lysosomal cobalamin 308 exporter ABCD4 where the two NBDs are in contact and the other from Salmonella typhimurium's lipid A transporter MsbA 309 with a widely open cavity. The overwhelming majority of conformations are regularly spaced by inter-NBD distance increments 310 smaller than 1 Å. By contrast, the sequence similarity-based collections populate sparse regions of this continuous transition, 311 with a high concentration of semi-open and open states (Fig. 3C, in pink and red, and Movie S2). 312

### 313 Classical manifold learning techniques can generate highly accurate conformations

Beyond describing the observed conformational variability, we evaluated the ability of two classical manifold learning 314 techniques, namely the linear PCA and the non-linear kernel PCA (kPCA), to generate unseen conformations. To do so, we 315 identified a set of ten conformational ensembles with very different degrees of motion complexity (Fig. 4A and Table S3). They 316 comprise between 20 and over 3 300 conformations and their reference chains contain 80 to 1 200 residues. They represent 317 proteins or protein families displaying substantial ( $\geq 5$  Å) and functionally relevant conformational changes, namely adenylate 318 kinase  $(ADK)^{62,63}$ , MurD<sup>19,64</sup>, the calcium pump ATPase<sup>65,66</sup>, the ABC transporters<sup>26,67</sup>, the small heat shock protein  $\alpha B$ 319 crystallin (Crys)<sup>68,69</sup>, the heat shock protein HSP90<sup>70,71</sup>, calmodulin (CALM)<sup>72,73</sup>, kinases (KIN)<sup>74,75</sup>, RAS<sup>25,76</sup>, and the 320 HIV capsid protein (CAP)<sup>77,78</sup>. Most of them have been extensively characterized by experimental structure determination 321 techniques or computational methods for simulating protein dynamics. Targeting their motions or their specific conformations 322 bears a therapeutic interest. 323

Within each ensemble, we first learned low-dimensional representations of a subset of conformations used as training 324 samples. We then projected the test conformations, not seen during training, to the learned representation space, and mapped 325 the projections back to the original 3D Cartesian space. The mapping is determined analytically in the case of linear PCA 326 and learned in the case of kPCA (see *Methods*). We evaluated the quality of the 3D reconstructions by computing their RMS 327 deviations from the original conformations. We found that both PCA and kPCA yield some high-accuracy reconstructions, with 328 an RMSD error below 1.5 Å, for all proteins (Fig. 4B). The error distribution width varies from one protein to another and does 329 not depend on motion complexity. For instance, all reconstructed conformations of HSP90 deviate by less than 2 Å from the 330 original ones, while the reconstruction error can be as high as 8 Å for MurD. The nonlinear kPCA performs significantly better 33



**Figure 3. ABC transporters' conformational variability. A.** Examples of protein structures from the ABC structure similarity-based conformational collection. The reference chain (5KOYA) is on the left, where we indicate the location of the two NBDs (~500 residues) and two TMDs (~700 residues). Within each of the other structures, we highlight one chain in marine, give its percentage of identity with the reference in squared brackets, and display the remaining chains in transparent grey. The six marine chains were assigned to six different collections by DANCE's default sequence similarity-based end-to-end protocol at  $l_{50}^{30}$ . **B.** Comparison of motion subspaces extracted from the sequence-based ensembles at  $l_{80}^{80}$  (61 conformations) and  $l_{50}^{30}$  (71 conformations) and the structure-based one (188 conformations). Each matrix shows the absolute pairwise scalar products computed for the first four PCA modes. The corresponding RMSIP are 0.99, 0.71 and 0.73. **C.** Distance between the geometric centres of the two NBDs (in Å). The conformations are ordered along the *x*-axis from the most closed one to the most open one.

than the linear PCA for all proteins from the benchmark. It allows increasing the percentage of high-quality reconstructions 332

(RMSD error <1.5Å) from 68 to 82% for MurD and from 1 to 33% for CALM (Table S4). Nevertheless, the reconstruction 333

accuracy of kPCA varies greatly depending on the values of the two hyperparameters controlling the kernel width and the 334

amount of regularisation (Fig. S6). The optimal values vary from one system to another and determining them *a priori* is not 335

trivial (**Table S5**). The applicability of non-linear techniques is thus limited by the choice of the adjustable parameters. 336

#### Reconstruction accuracy strongly depends on the distance to the training set 337

The quality of the predictions strongly correlates with the distance between the test conformation and the training set's convex 338 hull in the low-dimensional representation space (Fig. 4C). The linear PCA produces highly accurate reconstructions, with 339 an RMSD error smaller than 1.5 Å, only for conformations distant by less than 3 Å from the training set. We observed a 340 similar tendency for kPCA (Fig. S6). This dependence can be appreciated by visualising how the conformations cluster in the representation space (Fig. 4D). For instance, the most poorly reconstructed MurD conformation forms a singleton located far 342 away from all other conformations, particularly along the first most important principal component (Fig. 4D, dark dot). For this 343 protein, the kPCA performs substantially better than the PCA thanks to a better reconstruction of the most populated cluster Fig. **4D**, light squares). In addition, the overwhelming majority of conformations lie outside of the training set's convex hull. This 345 observation agrees with a recent study showing that interpolation almost surely never happens with high dimensional datasets<sup>79</sup> The 14 conformations located inside come from ADK, CALM, KIN, RAS and CAP and are all very well reconstructed, with 347

RMSD errors ranging from 0.1 to 1.7 Å. 348

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#### Influence of data uncertainty handling and conformation-specific centring 349

We assessed the influence of accounting for uncertainty in the data with position-wise weights and centring the data to a 350 reference conformation (Fig. S8-11). In principle, both operations may impact the conformations' superimposition and, as a 351 consequence, their final coordinates, as well as the extracted motions (see *Methods*). In practice, 95% of the ~35 000 ensembles 352 at  $I_{80}^{80}$  – excluding singletons and pairs, are not substantially altered by introducing position-wise uncertainty weights (Fig. S8). 353 They display the same displacement amplitude  $(\pm 1 \text{ Å})$  and motion complexity  $(\pm 1 \text{ mode})$ . When the weights are impactful, 354 they effectively lower the importance of large deviations in uncertain regions, *i.e.*, poorly covered by the conformations, and 355 prevent the associated motions, typically highly localised, from dominating the variance (Fig. S8, red dots). Hence, the 356 uncertainty weights tend to induce smaller deviations (Fig. S8A), increased motion complexities (Fig. S8B), and less dominant 357 and more collective main motions (Fig. S8C-D). 358 The choice of the reference conformation used for superimposing and centring the 3D coordinates has a much stronger 359 influence (Fig. S9). Only 43% of the  $l_{80}^{80}$  ensembles remain unaffected upon changing the reference. In this experiment, the

360 first reference is the multiple sequence alignment consensus (see *Methods*), while the second reference maximises the RMS 361 deviation from the first one. We expect this setup to yield the most contrasted resFig. S7ults. It almost never happened that 362 an ensemble consistently displayed a high motion complexity or a weakly contributing main motion for both references (Fig. 363 **S9B-C**). This result suggests that the ensembles exhibiting complex conformational rearrangements (e.g., loop deformations) 364 among a bulk of conformations also include a few conformations comparatively far from all the others. The motions simplify 365 when performing the PCA from the perspective of this minority. Normalising out the variance to focus on inter-residue 366 correlations attenuates this effect (Fig. S10). 367

#### Discussion 368

This work proposes a new perspective on the variability of protein 3D conformations. It provides the community with 369 conformational collections representing the multiple protein states available in the PDB and a fully automated versatile 370 computational pipeline to build custom collections. In doing so, it contributes to the representation and managing of multiple 371 conformational models of proteins. It enhances access and understanding of protein functional states and motions and facilitates 372 predictive methods benchmarking. Both DANCE pipeline and the produced PDB-wide data are readily usable in other studies. 373

We chose to rely on classical principal component analysis because of its intuitive geometrical interpretation. It allows 374 describing protein conformational variability with a limited set of orthogonal vectors interpretable as linear motions. We 375 provided estimates of motion complexity as the number of PCA components necessary to explain most of the observed 376 conformational variability. We found that a few linear motions suffice to explain most conformational collections. The high 377 complexity exhibited by a few protein families may reflect nonlinear structural deformations or seemingly random fluctuations. 378 For instance, protein kinases exhibit highly complex loop conformational rearrangements despite a well-conserved overall fold 379 and only two metastable functional states. Our analysis helps to identify such cases to prioritise their in-depth characterisation 380 with more sophisticated nonlinear dimensionality reduction techniques. 38.

We designed DANCE for dealing primarily with single polypeptidic chains grouped based on sequence similarity. To 382 go further, we have provided a proof-of-concept application study of DANCE's usefulness for comprehensively describing 383



**Figure 4.** Assessment of classical manifold learning techniques. A. Properties of the benchmark set. For each property *y*, we computed its normalised value as  $\frac{log(y) - \min(log(y))}{\max(log(y)) - \min(log(y))}$ . The minimum and maximum are determined over the whole  $l_{80}^{80}$  database. They are given at the bottom and on the top, respectively. **B.** Distributions of the RMSD reconstruction errors (in Å) for each ensemble in the benchmark set. We systematically reconstructed each conformation through a leave-one-cluster-out cross-validation procedure (see *Methods*). We set the two hyper-parameters of the kPCA to the values yielding the best reconstruction, for each ensemble. The protein names in the *x*-axis are ordered according to motion complexity. The stars indicate the statistical significance of the better performance of kPCA compared to linear PCA (one-sided paired t-test; \*: p-val <  $1e^{-2}$ ; \*\*\*: p-val <  $1e^{-5}$ ). **C.** RMSD reconstruction error in function of the distance to the training set's convex hull in the PCA representation space.

continuous motions shared across very distant homologs comprising different numbers of chains. We showed that ABC proteins
 with a wide diversity of substrates and transport mechanisms share a highly collective high amplitude opening/closing motion
 underlying their functioning. In addition, our work goes beyond a descriptive analysis by showing that classical manifold
 learning techniques can generate plausible conformations in the vicinity of the training set. Our results can serve as baselines
 for evaluating more sophisticated approaches.

DANCE superimposes the conformations onto representative references and describes conformational variability as a set of linear motions of these references. This approach offers a multi-view perspective on a given collection of conformations, easing interpretability and allowing for augmenting data in a learning context. Nevertheless, radical differences between conformations, such as fold changes, might confound the superimposition. Another limitation comes from the dependency of the superimposition on the multiple sequence alignment heuristic. Ambiguities arising from sequence similarities might result in suboptimal 3D coordinates matching and, thus, in large deviations. Future improvements will explore multi-reference or reference-free probabilistic frameworks and more refined accounts of data uncertainty<sup>80–84</sup>.

### **Data availability**

We provide public access to the conformational collections compiled by DANCE from the PDB at two levels of sequence similarity, namely  $l_{80}^{80}$  and  $l_{50}^{30}$  on Figshare. This repository also contains the structural similarity-based ABC transporter conformational collection along with the supplementary **Movies S1** and **S2**. In addition, we provide detailed information about the benchmark set and the assessment of PCA and kPCA.

### 401 Code availability

DANCE source codes are written in C/C++ and Python and are publicly available on GitHub at https://github.com/

<sup>403</sup> PhyloSofS-Team/DANCE. This repository also contains a Python wrapper allowing users to seamlessly run DANCE full

<sup>404</sup> pipeline. In addition, we provide example input 3D structures.

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### 572 Author contributions statement

<sup>573</sup> S.G. and E.L. designed research. V.L. and S.G. carried out the implementation. V.L., E.L. and S.G. produced and analysed the <sup>574</sup> results. E.L. wrote the manuscript with support and feedback from all authors. S.G. and E.L. supervised the project.

### **575** Competing interests

<sup>576</sup> The author(s) declare no competing interests.