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# Predicting protein–nucleic acid flexibility using persistent sheaf Laplacians

Nicole Hayes, <sup>a</sup> Ekaterina Merkurjev <sup>\*ab</sup> and Guo-Wei Wei <sup>acd</sup>

Understanding the flexibility of protein–nucleic acid complexes, often characterized by atomic *B*-factors, is essential for elucidating their structure, dynamics, and functions, such as reactivity and allosteric pathways. Traditional models such as Gaussian network models (GNM) and elastic network models (ENM) often fall short in capturing multiscale interactions, especially in large or complex biomolecular systems. In this work, we apply the Persistent Sheaf Laplacian (PSL) framework for the *B*-factor prediction of protein–nucleic acid complexes. The PSL model integrates multiscale analysis, algebraic topology, combinatorial Laplacians, and sheaf theory for data representation. It reveals topological invariants in its harmonic spectra and captures the homotopic shape evolution of data with its non-harmonic spectra. Its localization enables accurate *B*-factor predictions. We benchmark our method on three diverse datasets, including protein–RNA and nucleic-acid-only structures, and demonstrate that PSL consistently outperforms existing models such as GNM and multiscale FRI (mFRI), achieving up to a 21% improvement in Pearson correlation coefficient for *B*-factor prediction. These results highlight the robustness and adaptability of PSL in modeling complex biomolecular interactions and suggest its potential utility in broader applications such as mutation impact analysis and drug design.

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## 1 Introduction

Proteins and nucleic acids, including DNA and RNA, are some of the most essential building blocks of life. Proteins are involved in many vital processes, including cell signaling, gene regulation, transcription, and translation. Some of the key functions of proteins are binding (*e.g.*, to DNA in DNA polymerase), catalysis (*e.g.*, DNA polymerase catalyzing DNA replication), acting as molecular switches (*e.g.*, GTPases catalyzing GTP hydrolysis, in turn switching “off” cellular processes), and providing structure to cells and organisms (*e.g.*, actin, collagen, keratin, and silk).<sup>1</sup> Nucleic acids frequently act in conjunction with proteins to carry out important functions, such as gene expression and the storage and transmission of genetic information.<sup>2,3</sup> One of the most notable protein–nucleic acid complexes is the ribosome, mostly composed of RNA in addition to many smaller proteins, which synthesizes proteins and connects amino acids into polymer chains.<sup>3</sup> Another key complex is RNA polymerase, an enzyme that carries out

transcription, the process of synthesizing RNA using a DNA sequence, or template.<sup>1</sup>

Protein rigidity and flexibility are both crucial for protein structure and function.<sup>4,5</sup> Protein rigidity gives rise to the three-dimensional (3D) structure of proteins, and this 3D structure determines protein functions.<sup>1,6</sup> Protein flexibility plays a role in particular protein functions<sup>7</sup> like folding and interactions with other molecules, including nucleic acids. Nucleic acid flexibility is crucial for other biological processes, such as the role of DNA and RNA flexibility in packing as well as interactions between nucleic acids and proteins.<sup>3,8</sup> While there has been extensive study of protein flexibility in recent decades,<sup>9–12</sup> historically, much research on the flexibility of biomolecules primarily considered flexibility in the context of molecule motion and interaction. However, due to the discovery that proteins undergo thermal fluctuations even in neighborhoods of their native conformations (*i.e.*, folded states), flexibility is now understood to be an intrinsic property of proteins.<sup>13,14</sup>

There are multiple experimental approaches to measuring the flexibility of biomolecules, including X-ray crystallography, nuclear magnetic resonance (NMR), and single-molecule force experiments.<sup>15</sup> Protein flexibility, which can be measured by the *B*-factor, also called the Debye–Waller factor or temperature factor, is defined using the mean displacement of a scattering center during X-ray diffraction due to the thermal motion of atoms.<sup>16</sup> In addition to giving insight about the flexibility of atoms and amino acids in a protein structure, the *B*-factor also

<sup>a</sup> Department of Mathematics, Michigan State University, MI 48824, USA.  
E-mail: merkurje@msu.edu

<sup>b</sup> Department of Computational Mathematics, Science and Engineering Michigan State University, MI 48824, USA

<sup>c</sup> Department of Electrical and Computer Engineering, Michigan State University, MI 48824, USA

<sup>d</sup> Department of Biochemistry and Molecular Biology, Michigan State University, MI 48824, USA



provides information about other protein functions, including the protein's thermal motion, activity, and structural stability.<sup>17</sup>

Due to the significant differences in proteins and nucleic acids, models attempting to analyze the flexibility of protein–nucleic acid complexes must account for this variability. For instance, amino acid residues and nucleotides—the building blocks of proteins and nucleic acids, respectively—have different length scales, and thus multiscale models such as PSL are advantageous for capturing this information.<sup>3,18</sup> Many existing models for flexibility analysis utilize only one scale parameter, limiting their applicability on molecules with interactions over a wide range of scales. Notable examples include the anisotropic network model (ANM)<sup>19</sup> and Gaussian network model (GNM),<sup>20–22</sup> which are types of elastic network models (ENMs) and are some of the most popular methods for protein flexibility analysis.<sup>23–25</sup> ENMs treat the protein as a network, where the nodes are represented by the amino acid residues.<sup>19,21,26–29</sup> The first few eigenvalues of the network connectivity matrix reveal the long-time dynamics of proteins and can be used to predict *B*-factors. This process allows for the analysis of large proteins whose dynamics at large time scales would be intractable to traditional molecular dynamics (MD) simulations.<sup>23,30</sup>

The Gaussian network model typically outperforms the anisotropic network model in predicting *B*-factors,<sup>24,31</sup> and GNM has also been shown to be about one order more efficient than other existing flexibility analysis models.<sup>30</sup> Despite its advantages and popularity, GNM experiences difficulty in making predictions for many large biomolecules.<sup>32,33</sup> Park *et al.*<sup>31</sup> conducted extensive experiments applying GNM to predict *B*-factors on sets of relatively small, medium, and large protein structures. Although GNM achieved better results than normal mode analysis (NMA) on these data sets, it was unable to accurately predict *B*-factors for many structures.

Additional methods have emerged to overcome the disadvantages of GNM and ANM, including differential geometry analysis<sup>34</sup> as well as multiscale GNM (mGNM) and multiscale ANM (mANM) methods, which significantly improve protein *B*-factor prediction with respect to traditional GNM and ANM.<sup>25</sup> In particular, traditional GNM uses a single cutoff distance, limiting its predictive ability for molecules with interactions at multiple length scales, thereby motivating the mGNM and mANM methods. Another particularly successful method is the flexibility-rigidity index (FRI),<sup>23</sup> which is built on the theory of continuum elasticity with atomic rigidity (CEWAR). FRI is a structure-based approach that relies on two assumptions: that protein functions are entirely determined by a protein's structure and environment, and that protein structure is determined by a protein's interactions.<sup>3</sup> These assumptions allow FRI to bypass the Hamiltonian interaction matrix used in ENMs, leading to significantly lower computational complexity. Adaptations such as fast FRI (fFRI)<sup>24</sup> have further streamlined the process. Additionally, multiscale FRI (mFRI) has been shown to improve the performance of FRI on certain challenging protein structures for GNM, again largely due to the single cutoff distance employed by GNM.<sup>35</sup> Both GNM<sup>36</sup> and FRI<sup>3</sup> (including mFRI) have also been used to predict the flexibility of protein–

nucleic acid complexes, with two-kernel mFRI demonstrating marked improvement over both GNM and FRI.<sup>3</sup> More recently, an FRI method has been utilized for chromosome flexibility analysis, slightly improving predictive accuracy and significantly improving computational efficiency compared to GNM.<sup>37</sup>

Topological data analysis (TDA) has also been used as a technique for biomolecular study, and persistent homology has emerged as a particularly useful topological tool.<sup>38,39</sup> TDA has been widely applied to molecular sciences.<sup>40</sup> However, persistent homology has several drawbacks, including its inability to capture non-topological information.<sup>41</sup> To address these drawbacks, persistent topological Laplacians (PTLs), also simply called persistent Laplacians, were introduced by Wei and his coworkers in 2019<sup>41,42</sup> as a new approach to integrate the topological and geometric knowledge gained from persistent homology and multiscale graphs, respectively. Many other PTLs have since been developed, including the persistent path Laplacian,<sup>43</sup> persistent directed graph Laplacian,<sup>44</sup> persistent hyperdigraph Laplacian,<sup>45</sup> and persistent sheaf Laplacian (PSL).<sup>46</sup> Like many TDA methods, most of these algorithms are global, making them ill-suited for flexibility analysis, which requires information about individual atom sites in a biomolecule.<sup>47</sup> However, PSLs are capable of generating local information, thereby providing features for individual atoms in a protein and enabling the prediction of flexibility at specific sites. Other local persistent homology and persistent spectral methods have been used for RNA data analysis, including RNA flexibility prediction.<sup>48,49</sup> In addition to encoding local information, PSLs also retain the benefits of other persistent topological methods by capturing geometric and non-topological information. For more detail about recent advances in TDA, we refer the reader to ref. 50.

Recently, we have demonstrated the success of the persistent sheaf Laplacian (PSL) model in predicting protein *B*-factors.<sup>18</sup> While many existing topological approaches to molecular biology produce information about a molecule as a whole, the PSL model enables atom-specific feature generation, supporting its use for protein flexibility analysis. Additionally, due to its use of cellular sheaves, the PSL model allows for the inclusion of non-spatial information in addition to the topological information inherent to many TDA methods, which contributes to its predictive ability.

In the present work, we extend the application of the PSL model from proteins to protein–nucleic acid complexes. Other models, such as multiscale FRI (mFRI), have similarly demonstrated success in both protein and protein–nucleic acid flexibility analysis. Thus, the multiscale nature of the PSL model, also a feature of persistent homology methods and other persistent topological methods, supports this extension. Furthermore, the PSL method demonstrates marked improvement over GNM on a benchmark data set, achieving a 21% increase in the average Pearson correlation coefficient compared to GNM for one representative model. In addition to its improvement over GNM, the PSL model also performs well compared to a two-kernel mFRI method from the literature.<sup>3</sup>



One benchmark data set contains a subset of nucleic-acid-only structures, on which the PSL model also excelled compared to mFRI. This promising performance on varied structures demonstrates the strengths of the PSL model and suggests potential utility for other vital and complex molecules with interactions at multiple scales.

The remainder of this manuscript is organized as follows: Section 2 presents the results of the present work on data sets from the literature, Section 3 provides discussion of our results as well as a few case studies, and Section 4 reviews persistent sheaf Laplacian theory and describes the method of PSL feature generation used in our experiments. Specifically, Section 2.1 introduces the data sets used for benchmarking the PSL model. Section 2.2 reviews the parameters used in generating the PSL features for our model. Section 2.3 presents the results of the PSL model on two protein–nucleic acid data sets, including comparisons of the PSL method to existing results. Section 2.4 presents new results for a set of protein–RNA structures, Section 3.1 contains a few selected protein–nucleic acid complex case studies to illustrate our model's advantages.

## 2 Results

### 2.1 Data sets and coarse-grained models

Given the demonstrated success of the PSL model for protein flexibility analysis,<sup>18</sup> the present work extends this analysis to complexes consisting of both proteins and nucleic acids. Accordingly, the experiments in this section were performed on three data sets of protein–nucleic acid structures. The first data set, introduced by Yang *et al.*,<sup>36</sup> contains 64 structures, 19 of which consist of only nucleic acids (no amino acids).<sup>3</sup> The second data set of 203 high-resolution protein–nucleic acid complexes was introduced by Opron *et al.*<sup>3</sup> The third data set of specifically protein–RNA structures was previously used by Liu *et al.*<sup>51</sup> and Harini *et al.*<sup>52</sup> in analysis of binding affinity changes upon mutation. For PDB structures containing residues with alternate locations, we removed the lower-occupancy atomic coordinates in preprocessing, as outlined by Opron *et al.*<sup>3</sup>

In order to effectively perform flexibility analysis on these data sets, we must utilize coarse-grained representations of the structures. In our earlier paper on protein flexibility analysis using the PSL model,<sup>18</sup> we used a typical coarse-grained representation for proteins consisting of only the  $C_\alpha$  atom from each residue. Similarly, Yang *et al.*<sup>36</sup> proposed three coarse-grained representations for nucleic acids, denoted M1, M2, and M3.

The M1 model consists of backbone P atoms for nucleotides and  $C_\alpha$  atoms for proteins (*i.e.*, one atom per nucleotide). The M2 model includes the atoms from M1 as well as sugar O4' atoms (*i.e.*, two atoms per nucleotide). The M3 model contains the atoms from M1 in addition to sugar C4' atoms and base C2 atoms (*i.e.*, three atoms per nucleotide).<sup>3,36</sup> Our experiments in the present paper utilize these three coarse-grained representations, and we compare our results to those of Opron *et al.*<sup>3</sup> and Yang *et al.*<sup>36</sup> using these same representations. Fig. 1 provides a visualization of the three coarse-grained models, indicating which atoms are included in each representation.

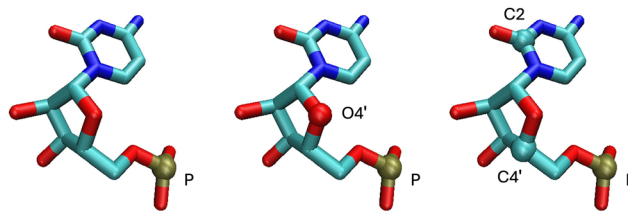


Fig. 1 Visualization of the three coarse-grained models used in this work using the visual molecular dynamics (VMD) software.<sup>53</sup> The figure illustrates models M1, M2, and M3, from left to right, for a single residue of nucleic acid 2TRA. Each representation depicts the atoms included in that model, and atoms are colored by name. The backbone P atom is included in each model, with models M2 and M3 including one and two additional atoms per nucleotide, respectively.

In addition to the data sets above, for which benchmark GNM results,<sup>36</sup> flexibility-rigidity index (FRI) results, and multiscale FRI (mFRI) results<sup>3</sup> exist in the literature, we also consider a data set of 141 protein–RNA complexes, which does not have existing *B*-factor prediction results. This data set was obtained from Liu *et al.*<sup>51</sup> and originally introduced by Harini *et al.*<sup>52</sup> Both of these studies concerned experiments on 710 mutations of these complexes. To generate a data set for our own experiments, we extracted the unique PDB IDs of the complexes from the data set of 710 mutations and performed our analysis using the PDB files of these complexes. (Note that in the original paper,<sup>52</sup> it is stated that the 710 mutations originate from 134 protein–RNA complexes. However, there are 141 unique PDB IDs in the data set of 710 mutations. Thus, the present work refers to the set of 141 protein–RNA complexes).

In our *B*-factor experiments, we excluded 15 of the 141 protein–RNA complexes due to unrealistic *B*-factors, resulting in a set of 126 structures. Specifically, PDB IDs 1AUD, 2JPP, 2KFY, 2KG0, 2KXN, 2LEB, 2LEC, 2LI8, 2M8D, 2MJH, 2MXY, 2RU3, 5M8I were excluded due to all atomic *B*-factors being equal to zero. The other two excluded structures were 2ERR (all *B*-factors were equal to 10.00) and 3J5S (all *B*-factors were equal to 1.00). We preprocessed the remaining 126 complexes as above to create M1, M2, and M3 models for each complex.

### 2.2 Parameters

For all sets of results in this work, we used filtration parameters of 6 Å, 12 Å, and 18 Å for the PSL model. Specifically, for each filtration radius, we constructed a 0th persistent sheaf Laplacian matrix  $L_0$  and computed its eigenvalues. To generate PSL features for each radius, we then computed the maximum, minimum, mean, and median of the nonzero eigenvalues of  $L_0$  and counted the number of its zero-valued eigenvalues. In addition to these five features for each of our three filtration radii, we also included the standard deviation of the nonzero eigenvalues of  $L_0$  for 18 Å, resulting in 16 total features for downstream regression tasks. For the experimental *B*-factor predictions in this section, we performed linear regression on each data set as a whole using the PSL features.



### 2.3 Results on 64 and 203 structures

In this section, we report our results for the PSL model for *B*-factor prediction of the protein–nucleic acid complexes in the data sets of 64 and 203 structures, respectively. The results for the data set of 64 complexes are displayed in Table 1, and the results for the 203-structure data set are shown in Table 3. Additionally, the results on nucleic-acid-only structures from the set of 64 complexes can be found in Table 2.

Our first set of results on the data set of 64 protein–nucleic acid complexes is shown in Table 1. We compare our findings to the flexibility-rigidity index (FRI) and multiscale FRI (mFRI) from Opron *et al.*<sup>3</sup> and the benchmark GNM.<sup>36</sup> The PSL model achieves a higher average Pearson correlation coefficient (PCC) than all other compared models for the M1, M2, and M3 representations. Most notably, our results using PSL features show an improvement over the GNM results by 21% for the M3 model.

In addition to the results on the entire data set, Opron *et al.*<sup>3</sup> also reported their average PCC for the subset of 19 nucleic-acid-only structures in the set of 64 complexes. Our results are compared to these in Table 2. Our PSL model achieved a higher average PCC value than that of the mFRI model for all three coarse-grained representations of the 19 structures in this set. In particular, we observe the biggest improvement for the M1 model, with our PSL model yielding an average PCC of 0.675, an 11% increase from the mFRI result (PCC = 0.608).

As shown in Table 3, our PSL model also achieved higher average PCC values on the set of 203 structures than the FRI and mFRI models.<sup>3</sup> Here, the most improvement is seen for the M3 representation, with the PSL model yielding an average PCC of 0.710, a 12% increase compared to mFRI (PCC = 0.63).

### 2.4 Results on 126 protein–RNA structures

In this section, we report our results for the prediction of *B*-factors of the 126 protein–RNA complexes described in Section 2.1. Table 4 shows the average PCC values over all 126 structures for the three models. The PSL model performed fairly consistently for all three models of this data set, with the M3 model yielding the highest average PCC (PCC = 0.700) of the three representations. Again, we utilized the same PSL features for these predictions as in our previous experiments, as outlined in Section 2.2. Of course, these parameters can be further tuned to optimize the results on the 126 protein–RNA

**Table 1** Average Pearson correlation coefficients (PCC) on three coarse-grained models for a set of 64 protein–nucleic acid structures.<sup>36</sup> Results are shown for our PSL model compared to the flexibility-rigidity index (FRI) and multiscale FRI (mFRI) models by Opron *et al.*<sup>3</sup> and the Gaussian network model (GNM).<sup>36</sup> The mFRI results shown were produced by a model using two kernels

Model	PSL	mFRI <sup>3</sup>	FRI <sup>3</sup>	GNM <sup>36</sup>
M1	<b>0.683</b>	0.666	0.620	0.59
M2	<b>0.669</b>	0.668	0.612	0.58
M3	<b>0.669</b>	0.620	0.555	0.55

**Table 2** Average Pearson correlation coefficients (PCC) for three coarse-grained models of a set of 19 nucleic-acid-only structures<sup>36</sup> using our PSL model. Results are compared to those of a multiscale flexibility-rigidity index (mFRI) model using two kernels<sup>3</sup>

Model	PSL	mFRI <sup>3</sup>
M1	<b>0.675</b>	0.608
M2	<b>0.649</b>	0.617
M3	<b>0.629</b>	0.603

**Table 3** Average Pearson correlation coefficients (PCC) for three coarse-grained models of a set of 203 protein–nucleic acid structures<sup>3</sup> using our PSL model. Results are compared to those of a flexibility-rigidity index (FRI) model and a multiscale flexibility-rigidity-index (mFRI) model using two kernels<sup>3</sup>

Model	PSL	mFRI <sup>3</sup>	FRI <sup>3</sup>
M1	<b>0.715</b>	0.68	0.613
M2	<b>0.718</b>	0.67	0.625
M3	<b>0.710</b>	0.63	0.586

**Table 4** Average Pearson correlation coefficients (PCC) for three coarse-grained models of a set of 126 protein–RNA structures<sup>52</sup> using our PSL model

Model	PSL
M1	0.669
M2	0.665
M3	0.700

structures in this data set. Detailed results for the 126 protein–RNA structures are given in Table 5 in Appendix A.

## 3 Discussion

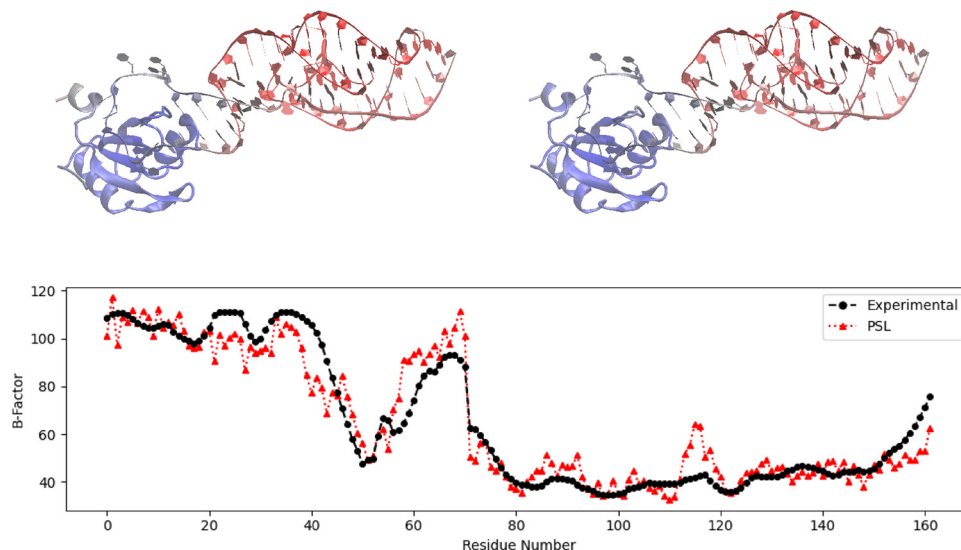
This section provides further discussion of the results presented in Section 2 and examines a few case studies of particular protein–nucleic acid complexes (and one nucleic-acid-only complex) to illustrate our model's strengths.

### 3.1 Protein–nucleic acid case studies

In addition to *B*-factor prediction results on the protein–nucleic acid data sets, in this section, we present case studies for selected protein–nucleic acid complexes to illustrate PSL model's advantage over mFRI on particular structures.

The first case study concerns complex PDB: 1DRZ, a hepatitis  $\delta$  virus ribozyme.<sup>36</sup> Ribozymes are RNA molecules (or protein–RNA complexes, such as complex 1DRZ) that can act as enzymes, catalyzing chemical reactions much like protein enzymes. However, the catalytic center of a ribozyme is made up solely of RNA, enabling catalysis without the need of a protein.<sup>54</sup> Fig. 2 displays the predicted *B*-factors for this complex using the PSL model compared to the experimental *B*-factors. Additionally, we provide a 3D visualization of the ribozyme using the Visual Molecular Dynamics (VMD)





**Fig. 2** Top: 3D visualization of complex 1DRZ, with residues colored by experimental  $B$ -factors (left) and predicted  $B$ -factors using PSL (right). Bottom: Experimental and predicted  $B$ -factors for each residue of complex 1DRZ. The results shown use model M1 for this complex. Both chains A and B of the complex are depicted here, with chain A corresponding to the atoms numbered 1–71 above and chain B corresponding to the atoms numbered 72–162 above. Note that the numbering used here simply reflects the ordering of atoms in the M1 model as extracted from the PDB file—this is not a canonical numbering for the complex.

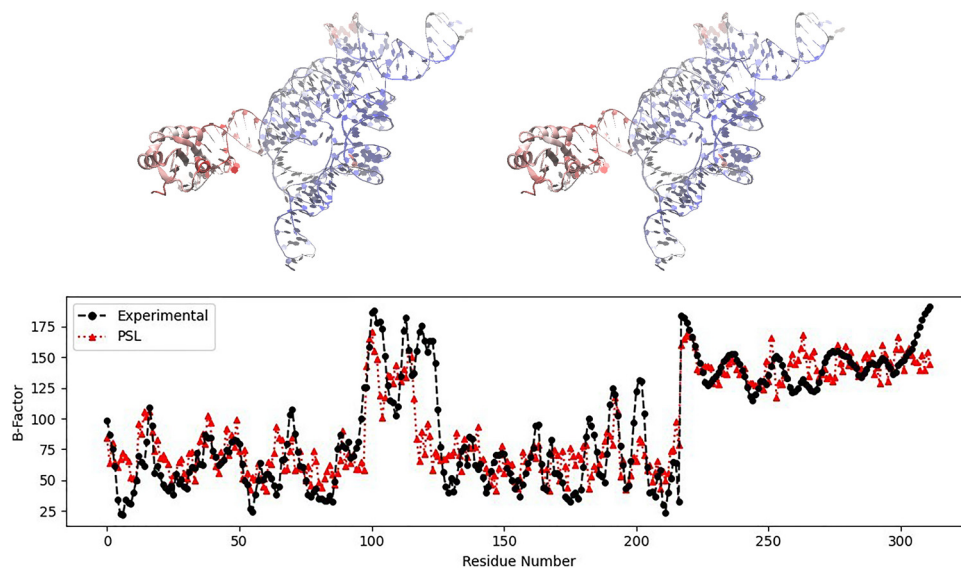
software.<sup>53</sup> Here, residues are colored by experimental or predicted  $B$ -factors. Less flexible regions are shown as blue (“colder” residues), and more flexible regions are shown as red (“warmer” residues). These predictions were generated using the M1 model for complex 1DRZ, which achieved the largest PCC value (PCC = 0.947) of all three coarse-grained models for this complex. The PSL model improves the result of the mFRI method (PCC = 0.846)<sup>3</sup> by 11%. Both the mFRI and PSL models performed best with the M1 coarse-grained model for this complex. Given our large PCC on this complex, the visualizations reflect this accuracy, with the PSL model slightly over- and undershooting the  $B$ -factors for a few sections of this ribozyme. For instance, for the short alpha helix corresponding to the last numbered residues, the PSL model predicted lower  $B$ -factors, reflected by this structure’s more saturated blue color in the 3D visualization.

Another protein–nucleic acid complex for which the PSL model outperforms the mFRI method is PDB: 1U6B, a Group I intron.<sup>36</sup> Group I introns are ribozymes that perform RNA self-splicing—that is, they catalyze reactions to excise themselves from the precursor RNA.<sup>55,56</sup> Fig. 3 shows the predicted and experimental  $B$ -factors for the complex 1U6B. The M1 model yielded the greatest PCC value of the three coarse-grained models on this complex as well, and the results shown are for this model. The PSL method achieved a PCC of 0.842, which is a 66% improvement over the mFRI model’s best result (PCC = 0.506) for this complex.<sup>3</sup> The mFRI model performed the best using the M3 model for this structure. The PSL model largely captures the variations in flexibility of complex 1U6B over all chains of the molecule, with some overestimations and underestimations at various areas, notably for one part of the flexible nucleic acid region corresponding roughly to residues 115–125 in Fig. 3 (in chain B). The PSL model successfully predicts the

increase in flexibility for the earlier residues of this flexible region but it underestimates the  $B$ -factors for the later residues of this region. Additionally, the  $B$ -factors for the flexible alpha helix corresponding to the last residues in Fig. 3 (in chain A) are somewhat underestimated by the PSL model. Still, the model is able to perform well overall, particularly compared to mFRI.

Our third case study concerns the nucleic-acid-only complex PDB: 2TRA, an aspartic acid transfer RNA (tRNA).<sup>36</sup> The primary function of tRNA lies in protein synthesis. Specifically, tRNA transports amino acids to the ribosome, where it then acts as a link between messenger RNA (mRNA) and the growing polypeptide chain. tRNA is also involved in other biological processes, including enzyme synthesis regulation, enzyme inhibition, and gene expression.<sup>57,58</sup> Fig. 4 presents the predicted  $B$ -factors for complex 2TRA using PSL as well as experimental  $B$ -factors. The results displayed are those for the M1 model, which achieved the highest PCC of the three coarse-grained models for this complex. Our PSL method yielded a PCC value of 0.744, a 21% improvement over mFRI (PCC = 0.614).<sup>3</sup> The mFRI result from Opron *et al.*<sup>3</sup> is also for the M1 model, which resulted in the highest PCC value for mFRI for this structure, with the M2 model performing equally well. Overall, the PSL model predicts the changes in flexibility throughout the nucleic acid well, while missing some of the finer points. The model successfully predicts the  $B$ -factors for the flexible residue numbered 46 in Fig. 4 and also captures the flexible region corresponding to the last numbered residues of the molecule for this model, while underestimating residue 62. While the PSL model does predict the other trends in flexibility, it misses some of the extremes—it underestimates the  $B$ -factors for some flexible regions (residues 0–4, 14–17) and overestimates the  $B$ -factors for some rigid regions (residues 7–11, 35–43).





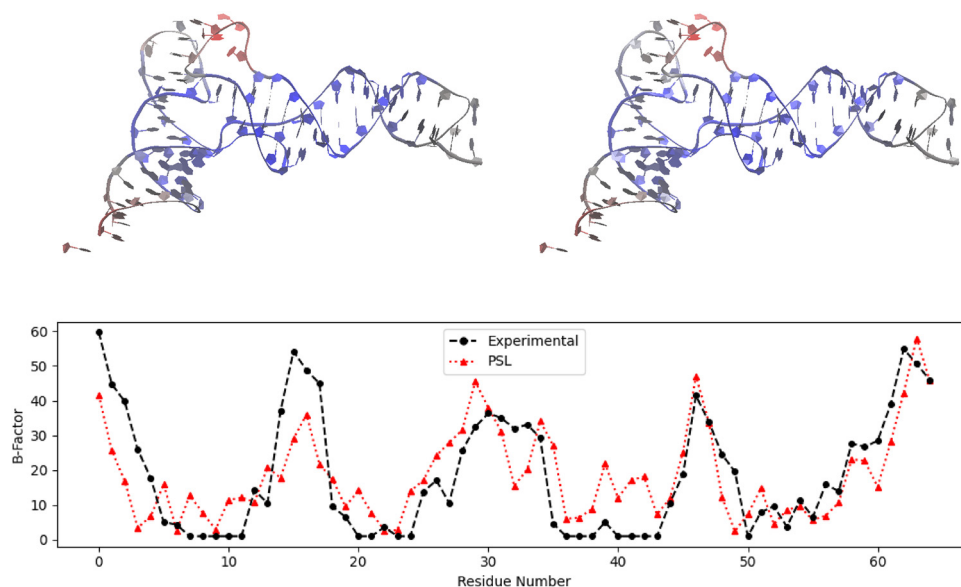
**Fig. 3** Top: 3D visualization of complex 1U6B, with residues colored by experimental  $B$ -factors (left) and predicted  $B$ -factors using PSL (right). Bottom: Experimental and predicted  $B$ -factors for each residue of complex 1U6B. The results shown use model M1 for this complex. Chains A, B, C, and D of the complex are depicted here, with chain A corresponding to the atoms numbered 218–312, chain B corresponding to the atoms numbered 0–195, and chain C corresponding to the atoms numbered 196–216. There is a single atom from Chain D numbered 217 in our model. Note that the numbering used here simply reflects the ordering of atoms in the M1 model as extracted from the PDB file—this is not a canonical numbering for the complex.

### 3.2 Discussion of results

Our results in Section 2 demonstrate the improved performance of the PSL model over traditional GNM and FRI as well as mFRI, a multiscale method. The successes of the PSL and mFRI models with respect to the single-scale methods illustrate the benefit of multiscale models for flexibility analysis of protein–nucleic acids, where interactions take place across multiple length scales. We have considered multiple data sets of protein–nucleic acid complexes containing diverse structures, including some nucleic-acid-only

complexes. Yang *et al.*<sup>36</sup> and Opron *et al.*<sup>3</sup> provide additional information about the complexes in the data sets of 64 and 203 structures.

In the set of 64 protein–nucleic acid complexes,<sup>36</sup> the coarse-grained molecule representations ranged from 61 atoms (for the M1 model representation for complex 1FIR) to 12 378 atoms (for the M3 model representation for complexes 1S72, 1YHQ, and 1YIJ). Overall, the PSL model performed the best on average using the M1 model for this data set (average PCC = 0.683). In fact, the M1 model yielded the highest PCC value for



**Fig. 4** Experimental and predicted  $B$ -factors for each residue of nucleic-acid-only complex 2TRA. The results shown use model M1 for this complex, which consists of only one chain.



all three complexes considered in our case studies in Section 3.1. These three complexes, 1DRZ, 1U6B, and 2TRA, were represented by models with a relatively small number of atoms: 162, 312, and 65, respectively, for the M1 model. These model sizes reflect the sizes of the complexes themselves—complex 1DRZ has 95 amino acid residues and 72 nucleotides, complex 1U6B has 95 residues and 222 nucleotides, and complex 2TRA has 74 nucleotides.<sup>36</sup> Thus, it is perhaps unsurprising that the PSL model using the M1 representation (with the fewest number of atoms per nucleotide) was sufficient to capture the topological and geometric information of these structures.

Generally, the PSL model performed better for smaller structures (*i.e.*, with fewer residues and/or nucleotides) than for larger structures for the 64-complex data set. This may be due to increased complexity of the larger molecules as well as the fact that we used the same filtration parameters for all complexes in the present work. Although our parameters may have generated features that encoded enough information about smaller molecules, adjusting the filtration radii to include greater length scales may improve the predictive performance of the PSL model on larger complexes by capturing potential longer-range interactions and additional structure around each atom. Additionally, although the M3 model had a lower PCC value on average for this data set, some of the larger molecules had better results with the M3 model than the M1 and M2 models. For instance, complex 1YIJ (a 50S ribosomal subunit with 3775 amino acid residues and 2876 nucleotides<sup>36</sup>) had a PCC of 0.498 using the M1 model (with 6636 atoms) and a PCC of 0.546 using the M2 model (with 9507 atoms), but using the M3 model (with 12 378 atoms) increased its PCC to 0.668. Consequently, we suggest that strategically choosing an appropriate coarse-grained model and tuning PSL filtration parameters based on molecular structure can improve flexibility prediction accuracy on a given complex.

The set of 203 structures<sup>3</sup> had many smaller molecules, with the complex 1VTG containing only seven atoms in its M1 representation. This data set also lacked such large molecules as the 64-structure data set—here, the complex with the most atoms in its coarse-grained representation was 4DQQ, with 1278 atoms using the M3 model. Given the above discussion, it is likely not surprising that the PSL model thus performed better on average for the 203-complex data set. This time, the M2 model yielded the best result (which was: average PCC = 0.718), although the M1 and M3 models showed very similar performance (which was: M1 average PCC = 0.715, M3 average PCC = 0.710). We do observe some similar trends for this data set as for the set of 64 complexes, with the PSL model generally achieving better predictive performance on the smaller structures.

In addition to the data sets of 64 and 203 protein–nucleic acid complexes, which have existing results in the literature for GNM, FRI, and mFRI, we also predicted *B*-factors for a data set of 126 protein–RNA complexes from a set of 710 mutations on these structures.<sup>51,52</sup> Detailed results and descriptions of the coarse-grained models for these 126 complexes are given in Appendix A. This data set did not contain structures as small as

those in the set of 203 complexes nor structures as large as those in the set of 64 complexes. For this data set, the PSL model performed best on average using the M3 representation (average PCC = 0.700). Furthermore, while the structures on which PSL performed the best were similar in size to the most successful complexes in the other two data sets, many of the larger structures in this data set had better results compared to similarly sized complexes in the other data sets.

Overall, the results of our PSL model on these three data sets demonstrate its applicability for flexibility analysis of diverse protein–nucleic acid complexes, as evidenced by its increased predictive accuracy over traditional GNM and mFRI, a state-of-the-art multiscale model. Our case studies further illustrate the advantages of PSL for particular significant structures. Additionally, we propose that the performance of the PSL model on larger complexes may be improved by parameter tuning and strategic coarse-grained model selection.

## 4 Methods

### 4.1 Persistent sheaf Laplacians

Many of the techniques used in the present work follow those in our prior paper,<sup>18</sup> with some adjustments made for the more complex nucleic acid structures. The algorithms for data preprocessing are described in Section 2.1, with particular discussion of three coarse-grained models from Yang *et al.*<sup>36</sup> that we adopted for our analyses. The specific persistent sheaf Laplacian features are also detailed in this paper in Section 2.2. In this section, we will briefly review the persistent sheaf Laplacian theory and provide more detail about the feature generation. The full details and comprehensive review of the persistent sheaf Laplacian theory and algorithms can be found in the original paper by Wei and Wei<sup>46</sup> and our earlier paper.<sup>18</sup> Other graph-based approaches incorporating Laplacians include the work.<sup>59</sup>

Persistent sheaf Laplacians<sup>46</sup> (PSLs) extend the theory of persistent Laplacians on simplicial complexes and simplicial chain complexes to cellular sheaves and sheaf cochain complexes. The motivation for the extension to sheaves is, in part, their ability to embed additional non-spatial information in a simplicial complex. In molecular science, this information may be derived from molecular structures; one example of such information is an atom's partial charge. In contrast to many other persistent topological Laplacians, which can describe a molecule as a whole, persistent sheaf Laplacians provide information about local (*i.e.*, atom-specific) topology and geometry in a molecule.

To formalize the above, we will first introduce persistent homology, which is defined with respect to simplicial complexes. Given a finite set  $V$  (for example, a set of atoms in a molecule), we can define a simplicial complex  $X$  on  $V$  as a collection of subsets of  $V$  satisfying the following criterion: if a set  $\sigma$  is in  $X$ , then any subset of  $\sigma$  must also be in  $X$ . The sets  $\sigma$  are called simplices, and a simplex with  $q + 1$  elements is called a  $q$ -simplex. Furthermore, if a simplex  $\sigma \subset \tau$ , then  $\sigma$  is called a face of  $\tau$ , and this face relation is defined by  $\sigma \leq \tau$ . We can



define subcomplexes as follows: if  $X$  and  $Y$  are both simplicial complexes with  $X \subset Y$ , then  $X$  is considered a subcomplex of  $Y$ .

Given a simplicial complex  $X$ , one may define the simplicial chain complex associated with  $X$ , which is denoted as

$$\cdots \xrightarrow{\partial_3} C_2(X) \xrightarrow{\partial_2} C_1(X) \xrightarrow{\partial_1} C_0(X) \rightarrow 0.$$

$C_q(X)$  is a real vector space generated by  $q$ -simplices of  $X$ , and an element  $\sigma \in C_q(X)$  is called a  $q$ -chain. The operator  $\partial_q: C_q(X) \rightarrow C_{q-1}(X)$  is a linear map, called a boundary map, defined as follows:

$$\partial_q[v_{a_0}, \dots, v_{a_q}] = \sum_i (-1)^i [v_{a_0}, \dots, \hat{v}_{a_i}, \dots, v_{a_q}].$$

Here, the notation  $\hat{v}_{a_i}$  means that the element  $v_{a_i}$  is deleted. Because our finite set  $V$  is totally ordered, the boundary operator  $\partial_q$  is well defined. We can then define the  $q$ -th homology group of the chain complex by  $H_q = \ker \partial_q / \text{im} \partial_{q+1}$ . We have  $\partial^2 = \partial_q \circ \partial_{q+1} = 0$ , so  $H_q$  is also well defined.

Now, if we let  $X$  be a subcomplex of  $Y$ , we also have inclusions of the chain groups, with the inclusion map denoted by  $\iota: C_q(X) \rightarrow C_q(Y)$ . Thus, we can construct the following diagram:

$$\begin{array}{ccccccc} \cdots & \xrightarrow{\partial_{q+2}^X} & C_{q+1}(X) & \xrightarrow{\partial_{q+1}^X} & C_q(X) & \xrightarrow{\partial_q^X} & C_{q-1}(X) & \xrightarrow{\partial_{q-1}^X} & \cdots \\ & & \downarrow & & \downarrow & & \downarrow & & \\ \cdots & \xrightarrow{\partial_{q+2}^Y} & C_{q+1}(Y) & \xrightarrow{\partial_{q+1}^Y} & C_q(Y) & \xrightarrow{\partial_q^Y} & C_{q-1}(Y) & \xrightarrow{\partial_{q-1}^Y} & \cdots \end{array}$$

Then, we have a map  $\iota^*$  induced by the inclusion  $\iota$ , defined as  $\iota^*: H_q(X) \rightarrow H_q(Y)$ . The image

$$\iota^*(H_q(X))$$

is the  $q$ -th persistent homology group for the simplicial complex pair  $(X, Y)$ . The ranks of these persistent homology groups are called ( $q$ -th) persistent Betti numbers  $\beta_q^{X,Y}$ ,<sup>46</sup> which are used in topological data analysis to track topological features that persist in the filtration (*i.e.*, at multiple scales). If the input point cloud is given by a set of atoms in a molecule, the Betti numbers can capture multiscale information about the molecule's structure and interactions.

Persistent Laplacians<sup>41</sup>—in particular, their non-zero eigenvalues—can capture even more information from an input point cloud. Given a persistent homology group  $\iota^*(H_q(X))$  as defined above, a persistent Laplacian is a positive semi-definite operator with a kernel isomorphic to that group. To construct the persistent Laplacian, first define  $C_{q+1}^{X,Y} = \{c \in C_{q+1}(Y) | \partial_{q+1}^Y(c) \in C_q(X)\}$ , and let  $\partial_{q+1}^{X,Y}$  be the restriction of  $\partial_{q+1}^Y$  to  $C_{q+1}^{X,Y}$ . Note that each  $C_q(X)$  is generated by  $q$ -simplices, equipping it with a canonical inner product. Now, we can define the  $q$ -th persistent Laplacian  $\Delta_q^{X,Y}$  as

$$\Delta_q^{X,Y} = \partial_{q+1}^{X,Y} (\partial_{q+1}^{X,Y})^\dagger + (\partial_q^X)^\dagger \partial_q^X,$$

where  $\dagger$  is the adjoint of a linear morphism. Many variants of persistent Laplacian methods exist, and examining the

eigenvalues of these various persistent Laplacians enables regression algorithms to learn multiscale information about point clouds in addition to that available from persistent homology.

Now, while persistent Laplacians as defined above are built on simplicial chain complexes, persistent sheaf Laplacian theory<sup>46</sup> generalizes this idea to cellular sheaves and sheaf cochain complexes. A cellular sheaf  $\mathcal{F}$  can be conceptualized as a simplicial complex  $X$ , where we assign (i) a finite-dimensional vector space  $\mathcal{S}(\sigma)$  to each simplex  $\sigma$  of  $X$  and (ii) a linear morphism  $\mathcal{S}_{\sigma \leq \tau}$  of vector spaces to each face relation  $\sigma \leq \tau$ . The vector space  $\mathcal{S}(\sigma)$  is called the stalk of  $\mathcal{S}$  over  $\sigma$ , and the morphism  $\mathcal{S}_{\sigma \leq \tau}$  is called the restriction map of the face relation  $\sigma \leq \tau$ . Furthermore, the restriction maps behave nicely across face relations; that is,  $\mathcal{S}(\sigma)$  satisfies the rule that if  $\rho \leq \sigma \leq \tau$ , then  $\mathcal{S}_{\rho \leq \tau} = \mathcal{S}_{\sigma \leq \tau} \mathcal{S}_{\rho \leq \sigma}$ , where  $\mathcal{S}_{\sigma \leq \sigma}$  refers to the identity map of  $\mathcal{S}(\sigma)$ .

Intuitively, stalks serve as information about each simplex; in the case where the input point cloud represents atoms in a molecule, a stalk may capture non-spatial atomic information, such as partial charges, atomic weights, *etc.* We can view restriction maps as describing how the additional information stored by stalks interacts across simplexes.

Now, we may construct a sheaf cochain complex

$$0 \rightarrow C^0(X; \mathcal{S}) \xrightarrow{d} C^1(X; \mathcal{S}) \xrightarrow{d} C^2(X; \mathcal{S}) \xrightarrow{d} \cdots$$

where each  $q$ -th sheaf cochain group  $C^q(X; \mathcal{S})$  is defined as the direct sum of stalks over all  $q$ -dimensional simplices. The coboundary maps  $d$  require obtaining a signed incidence relation by globally orienting the simplicial complex  $X$ , where the relation assigns an integer  $[\sigma: \tau]$  to each face relation  $\sigma \leq \tau$ . Thus, we can define the coboundary map  $d^q: C^q(X; \mathcal{S}) \rightarrow C^{q+1}(X; \mathcal{S})$  as

$$d^q|_{\mathcal{S}(\sigma)} = \sum_{\sigma \leq \tau} [\sigma: \tau] \mathcal{S}_{\sigma \leq \tau}.$$

To define the persistent sheaf Laplacian, we must again consider subcomplexes  $X \subset Y$ , now equipped with sheaf  $\mathcal{F}$  on  $X$  and sheaf  $\mathcal{G}$  on  $Y$ . Here, we let the sheaves be defined such that the stalks and restriction maps of  $X$  and  $Y$  are identical. Thus, we have a sheaf cochain complex for  $X$  and one for  $Y$ , with inclusion maps between their corresponding cochain groups. Further assuming that each stalk is an inner product space, we can define  $\Theta_{\mathcal{F}, \mathcal{G}}^{q+1} = \{x \in C^{q+1}(Y; \mathcal{G}) | (d_{\mathcal{G}}^q)^\dagger(x) \in C^q(X; \mathcal{F})\}$ , where  $d_{\mathcal{F}, \mathcal{G}}^q$  is the adjoint of  $\pi(d_{\mathcal{G}}^q)^\dagger|_{\Theta_{\mathcal{F}, \mathcal{G}}^{q+1}}: \Theta_{\mathcal{F}, \mathcal{G}}^{q+1} \rightarrow C^q(X; \mathcal{F})$  and  $\pi$  is the projection map from  $C^q(Y; \mathcal{G})$  to  $C^q(X; \mathcal{F})$  (its subspace). The  $q$ -th persistent sheaf Laplacian  $\Delta_q^{\mathcal{F}, \mathcal{G}}$  is then defined as

$$\Delta_q^{\mathcal{F}, \mathcal{G}} = (d_{\mathcal{F}, \mathcal{G}}^q)^\dagger d_{\mathcal{F}, \mathcal{G}}^q + d_{\mathcal{F}}^{q-1} (d_{\mathcal{F}}^{q-1})^\dagger.$$

Note that when  $\mathcal{F} = \mathcal{G}$ , the persistent sheaf Laplacian becomes the sheaf Laplacian of  $\mathcal{F}$ . Also, when  $\mathcal{F}$  and  $\mathcal{G}$  are constant, the persistent sheaf Laplacian is equal to the persistent Laplacian  $\Delta_q^{X,Y}$  defined above.



## 4.2 Model construction and feature generation

Given the theoretical foundation of PSLs in the previous section, in this section we will outline the specifics of the PSL model used in this paper, as well as how we generated spectral features using this model.

In general, persistent homology examines how topological features evolve across a nested sequence of subcomplexes, called a filtration. Given a point cloud, a common method for constructing a filtration is by varying a radius parameter from each point and including additional simplices once the radius parameter exceeds the distance from our point to each of those simplices. In the present work, the filtration parameter is the distance (in Å) from each atom. At each radius, one may compute topological invariants of the corresponding subcomplex. Then, how these invariants change over the filtration can be analyzed. In a molecule, this can allow the examination of interactions and structures at multiple length scales. However, many of the typical topological tools provide global information, whereas protein–nucleic acid flexibility analysis requires the knowledge of atom-specific information to predict individual  $B$ -factors. Persistent sheaf Laplacians provide the necessary localization for this analysis. To construct a filtration of sheaves, as used in persistent sheaf cohomology<sup>60</sup> and persistent sheaf Laplacians, we begin by constructing a filtration of simplicial complexes as above, but additionally construct a sheaf for each complex. Then, by computing the eigenvalues of persistent sheaf Laplacians, we can obtain topological invariants (from the harmonic spectra) and geometric information (from the non-harmonic spectra) of the data.<sup>45</sup>

Specifically, for the present work, we begin with a point cloud consisting of the atoms in a particular coarse-grained model (M1, M2, or M3) of a protein–nucleic acid complex. With the goal of predicting the  $B$ -factor for each atom, we construct features for each atom using PSLs. For a given atom  $A$  in the point cloud, we first designate a cutoff distance for the point cloud so that we only consider other atoms within that distance from  $A$  when we construct our complex. Then, we determine the set of radii that will generate our filtration—for this work, we used 6 Å, 12 Å, and 18 Å. For each radius, we construct an alpha complex  $X$  (again, only considering atoms within the prescribed cutoff distance from  $A$ ). To construct a cellular sheaf on  $X$ , we assign a label  $q_i$  to each atom  $v_i$  in  $X$ , and then let each stalk be. To define the restriction maps, let  $r_{ij}$  be the length of simplex  $v_i v_j$  (i.e., the 1-simplex between atoms  $v_i$  and  $v_j$ ). For face relations of the form  $v_i \leq v_i v_j$ , we define the restriction map as scalar multiplication by  $q_j/r_{ij}$ . For face relations of the form  $v_i v_j \leq v_i v_j v_k$ , we define the restriction map as scalar multiplication by  $q_k/(r_{ik}r_{jk})$ . (Further motivation behind the definition of restriction maps in this way can be found in our previous paper<sup>18</sup> and the introduction by Wei *et al.*<sup>46</sup>) In order to distinguish atom  $A$  from the other model atoms in  $X$ , the label  $q_i$  of  $A$  is set to 0, and the labels of all other model atoms in  $X$  are set to 1. By constructing sheaf Laplacians for atom  $A$  for a given radius, and then computing the eigenvalues of the sheaf Laplacians, we can generate features for  $A$  using these eigenvalues (the particular set of features used in this work is

detailed in Section 2.2). Then, by varying the radius and computing sheaf Laplacians across the filtration, we obtain additional features for each radius. We can repeat this process for all atoms in the given coarse-grained model to enable the  $B$ -factor prediction of each atom.

Regarding computational complexity, both GNM and the persistent sheaf Laplacian model used in this work have a computational complexity of  $O(n^3)$ , where  $n$  is the number of input nodes. The complexity of FRI and mFRI is  $O(n^2)$ , and this decreased complexity is one of the primary advantages of the FRI methods over GNM. Although the PSL model has greater computational complexity than the FRI models, the size of the present  $B$ -factor prediction is relatively small due to the truncation in our algorithm. In particular, when constructing a sheaf around each atom in a protein–nucleic acid complex, we do not consider all other atoms in the structure, but only a small subset of atoms within a chosen cutoff distance. Therefore, computing  $B$ -factors with the proposed method is quite manageable given its lower cost in practice.

## 5 Conclusion

Persistent topological Laplacians are a new generation of multiscale spectral algorithms.<sup>50</sup> Unlike other topological techniques, the persistent sheaf Laplacian (PSL) offers atom-specific localized information of a biomolecular complex.<sup>46</sup> The atom-specific and multiscale nature of persistent sheaf Laplacian features enables detailed atomistic analysis, which is crucial for understanding multiscale molecular flexibility and long-term conformational dynamics. Both molecular flexibility and the long-term conformational dynamics are very important for their biological functions and are the subjects of biophysical studies in the past decades.

In this study, we introduce a PSL model to predict the  $B$ -factors of protein–nucleic acid complexes, a task that presents unique challenges due to their multiscale nature and structural diversity. Our results across multiple benchmark datasets demonstrate that PSL not only surpasses traditional models like GNM and mFRI in predictive accuracy but also maintains strong performance on nucleic-acid-only structures. Furthermore, we have given a few case studies to further illustrate the model's effectiveness in capturing biologically relevant flexibility patterns. The promising performance of our PSL model suggests its potential for further applications in structural biology, including the prediction of mutation effects and binding affinity changes as well as the design of biomolecular therapeutics. To be consistent with the literature, a regression model is used in this work. However, the proposed sheaf approach can be implemented in a deep neural network for  $B$ -factor predictions.<sup>18</sup>

## Author contributions

Nicole Hayes was responsible for the software, computational experiments and writing. Ekaterina Merkurjev and Guo-Wei Wei were responsible for methodology, reviewing, editing, some writing and supervision.



## Conflicts of interest

The authors have no conflicts to disclose.

## Data availability

The code for the PSL model, including an example of PSL feature generation and *B*-factor prediction is available at [https://github.com/weixiaoqimath/persistent\\_sheaf\\_Laplacians](https://github.com/weixiaoqimath/persistent_sheaf_Laplacians).

## Appendix

### A Appendix: additional tables

Table 5.

**Table 5** Pearson correlation coefficients (PCC) for our *B*-factor predictions of 126 protein–RNA structures<sup>52</sup> using PSL features. Results are given for the three coarse-grained models M1, M2, and M3 of each structure, and the number of atoms in each model is also included

PDB ID	M1 Model		M2 Model		M3 Model	
	PCC	# of Atoms	PCC	# of Atoms	PCC	# of Atoms
1ASY	0.647	1114	0.651	1248	0.657	1382
1C9S	0.785	1583	0.804	1638	0.718	1693
1DFU	0.69	130	0.744	168	0.805	206
1FEU	0.478	450	0.528	530	0.609	610
1GTF	0.594	1571	0.577	1615	0.567	1659
1JBR	0.71	356	0.695	417	0.706	478
1JBS	0.637	332	0.584	366	0.552	400
1K8W	0.547	324	0.544	345	0.562	366
1QFQ	0.774	50	0.59	65	0.717	80
1SI2	0.843	128	0.719	137	0.721	146
1TTT	0.628	1401	0.624	1587	0.57	1773
1U0B	0.798	535	0.845	609	0.814	683
1URN	0.813	345	0.839	401	0.831	456
1UTD	0.515	1593	0.544	1628	0.516	1663
1WNE	0.62	487	0.604	500	0.575	513
1YTU	0.687	836	0.721	855	0.741	874
1YVP	0.671	1101	0.657	1145	0.648	1189
1ZDI	0.718	415	0.81	444	0.712	473
1ZH5	0.637	377	0.624	395	0.537	413
2BX2	0.329	514	0.39	528	0.454	542
2C4R	0.67	501	0.653	511	0.665	521
2E9T	0.524	974	0.597	1002	0.555	1030
2G4B	0.595	179	0.66	186	0.693	193
2I91	0.624	1087	0.662	1132	0.65	1177
2IX1	0.645	656	0.625	669	0.582	682
2LA5	0.675	52	0.388	88	0.289	124
2PJP	0.758	144	0.794	167	0.793	190
2X1A	0.711	88	0.75	89	0.689	90
2XFM	0.836	126	0.822	133	0.777	140
2XGJ	0.541	1750	0.54	1760	0.536	1770
2XS2	0.704	93	0.704	99	0.82	105
2Y8W	0.664	234	0.725	254	0.779	274
2YH1	0.65	203	0.629	212	0.584	221
2Z10	0.796	153	0.833	191	0.816	229
2ZKO	0.9	182	0.792	224	0.867	266
2ZZM	0.693	413	0.698	497	0.791	581
2ZZN	0.486	813	0.49	955	0.497	1097
3AM1	0.587	317	0.516	398	0.549	479
3EQT	0.409	283	0.427	299	0.466	315
3GIB	0.743	200	0.734	209	0.685	218
3GPQ	0.545	622	0.561	626	0.572	629
3K49	0.772	1086	0.759	1116	0.731	1146
3K5Q	0.73	409	0.723	418	0.716	427

Table 5 (continued)

PDB ID	M1 Model		M2 Model		M3 Model	
	PCC	# of Atoms	PCC	# of Atoms	PCC	# of Atoms
3K5Y	0.725	408	0.69	417	0.663	426
3L25	0.78	506	0.76	522	0.785	538
3MOJ	0.678	144	0.682	213	0.718	282
3NCU	0.818	262	0.862	284	0.87	306
3OL6	0.633	1984	0.627	2124	0.669	2264
3QGB	0.701	408	0.673	417	0.667	426
3QGC	0.665	408	0.651	417	0.644	426
3QSU	0.651	859	0.636	867	0.645	875
3RW6	0.654	609	0.575	729	0.688	849
3SN2	0.562	876	0.571	905	0.544	934
3U4M	0.462	308	0.598	388	0.718	468
3UZS	0.607	1032	0.635	1052	0.649	1072
3V71	0.829	371	0.813	378	0.829	385
3V74	0.584	405	0.571	416	0.566	427
3VYX	0.666	115	0.648	135	0.655	155
3VYY	0.861	215	0.896	265	0.91	315
3WBM	0.646	393	0.656	443	0.679	493
3ZLA	0.676	1919	0.671	2007	0.616	2095
4CIO	0.954	103	0.959	110	0.959	117
4ED5	0.537	355	0.548	373	0.535	389
4ERD	0.639	256	0.556	300	0.583	344
4HT8	0.743	374	0.736	388	0.715	400
4HT9	0.717	190	0.696	199	0.692	208
4M59	0.491	1400	0.513	1432	0.506	1463
4MDX	0.723	238	0.686	247	0.65	256
4NGB	0.695	287	0.698	299	0.686	311
4NGD	0.571	292	0.538	304	0.599	316
4NHA	0.682	246	0.732	262	0.757	278
4NKU	0.65	632	0.653	636	0.649	640
4O26	0.795	562	0.785	656	0.68	750
4OE1	0.492	1400	0.509	1432	0.501	1463
4OIO	0.749	397	0.763	399	0.765	401
4OOG	0.615	514	0.644	548	0.702	582
4PMW	0.595	1410	0.595	1438	0.576	1466
4QEI	0.638	630	0.701	698	0.761	766
4QG3	0.412	308	0.457	388	0.533	468
4QVC	0.745	367	0.741	370	0.739	373
4QVD	0.767	367	0.723	371	0.704	375
4QVI	0.41	309	0.481	389	0.585	469
4R3I	0.753	167	0.747	171	0.744	175
4R8I	0.803	68	0.803	68	0.803	68
4RCJ	0.798	191	0.782	194	0.755	197
4TUW	0.76	193	0.591	247	0.538	299
4V4F	0.732	221	0.599	228	0.675	235
4YVI	0.604	559	0.685	630	0.759	701
4Z4C	0.654	830	0.656	857	0.696	884
4Z4D	0.602	831	0.605	859	0.619	886
5AWH	0.689	1544	0.668	1616	0.682	1688
5DET	0.675	193	0.717	202	0.694	211
5DNO	0.785	170	0.639	177	0.639	184
5E3H	0.532	665	0.549	689	0.605	713
5EEU	0.757	1565	0.774	1609	0.682	1653
5EIM	0.723	332	0.69	348	0.688	364
5ELH	0.652	286	0.612	291	0.64	296
5ELK	0.675	126	0.682	130	0.678	134
5EN1	0.801	190	0.826	197	0.826	204
5EV1	0.735	204	0.731	211	0.692	218
5F98	0.55	4021	0.591	4165	0.634	4309
5GXH	0.728	678	0.715	684	0.706	690
5H1K	0.697	1388	0.686	1413	0.683	1438
5H1L	0.816	688	0.81	695	0.807	702
5HO4	0.726	188	0.705	198	0.712	206
5IP2	0.668	702	0.655	713	0.657	724
5JBJ	0.514	651	0.522	675	0.548	699
5M3H	0.65	2221	0.581	2249	0.664	2277
5MPG	0.434	103	0.429	110	0.418	117
5MPL	0.362	107	0.333	113	0.379	119
5UDZ	0.715	323	0.658	370	0.653	416
5W1H	0.72	1342	0.713	1383	0.716	1424



Table 5 (continued)

PDB ID	M1 Model		M2 Model		M3 Model	
	PCC	# of Atoms	PCC	# of Atoms	PCC	# of Atoms
5WLH	0.708	1332	0.705	1374	0.719	1416
5WWW	0.712	99	0.645	105	0.555	111
5WWX	0.644	89	0.663	94	0.671	99
5WZH	0.671	543	0.664	554	0.686	565
5YTV	0.849	79	0.852	83	0.821	87
5ZTM	0.613	380	0.633	428	0.665	476
6B14	0.797	523	0.785	606	0.667	689
6CMN	0.611	117	0.708	144	0.738	171
6CYT	0.715	669	0.759	686	0.773	703
6D12	0.928	237	0.949	275	0.93	313
6KWR	0.801	487	0.848	521	0.844	555
6NY5	0.721	769	0.712	782	0.725	795
6SDY	0.325	107	0.228	141	0.323	175
6SO9	0.735	117	0.742	122	0.752	127

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