

Hydrodynamic Radii of Intrinsically Disordered Proteins: Fast Prediction by Minimum Dissipation Approximation and Experimental Validation

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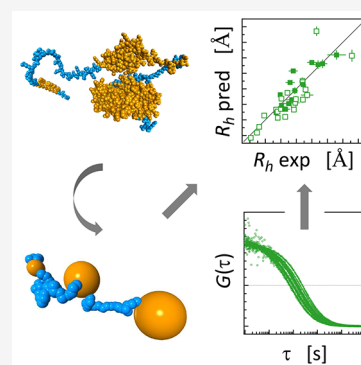
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ABSTRACT: The diffusion coefficients of globular and fully unfolded proteins can be predicted with high accuracy solely from their mass or chain length. However, this approach fails for intrinsically disordered proteins (IDPs) containing structural domains. We propose a rapid predictive methodology for estimating the diffusion coefficients of IDPs. The methodology uses accelerated conformational sampling based on self-avoiding random walks and includes hydrodynamic interactions between coarse-grained protein subunits, modeled using the generalized Rotne–Prager–Yamakawa approximation. To estimate the hydrodynamic radius, we rely on the minimum dissipation approximation recently introduced by Cichocki et al. Using a large set of experimentally measured hydrodynamic radii of IDPs over a wide range of chain lengths and domain contributions, we demonstrate that our predictions are more accurate than the Kirkwood approximation and phenomenological approaches. Our technique may prove to be valuable in predicting the hydrodynamic properties of both fully unstructured and multidomain disordered proteins.



Intrinsically disordered proteins (IDPs) constitute an extensive class of biological macromolecules, and their role in the homeostasis of a living cell has been increasingly recognized in recent decades.^{1,2} The frequency of long intrinsically disordered regions (IDRs) in proteins differs significantly between the kingdoms of life, ranging from 2% in archaea to 33% in eukaryotes.³ The IDP molecules display different degrees of structural disorder. Their chains can encompass either several folded globular domains or super-secondary structures connected by flexible linkers, sparse secondary structural elements, or can be completely natively unstructured. Disordered proteins exhibit a notable characteristic, the absence of a stable, well-defined relative spatial arrangement of their fragments. Instead, their equilibrium properties can be described through a broad set of rapidly interconverting conformers, posing a challenge for analysis, particularly in the context of long chains.⁴

The average geometric properties of IDPs, including their shape and size, are determined by the equilibrium ensemble of conformational states. This equilibrium state is intricately influenced by environmental conditions,⁵ such as temperature,⁶ ionic strength,^{7,8} osmolality,⁹ crowding,¹⁰ post-translational modifications,¹¹ and the presence of specific molecular binding partners.¹² The formation of transient or more stable non-covalent complexes introduces another nontrivial dependence of the IDP equilibrium geometry on environmental factors.

Because the shape and availability of the binding sites necessary for the interaction of IDP with ligands, other proteins, and nucleic acids are strongly influenced by the environment, IDPs often act as higher-order regulators in key cellular processes such as gene expression,^{11,13} signaling,^{2,14} or extracellular biomineralization.¹⁵ The different conformations of these flexible proteins enable IDPs to perform their multiple functions.¹ In particular, it is worth emphasizing the important roles of IDPs in health and disease, e.g., the role of the p53 protein as a tumor suppressor,¹⁶ mutations of which are often responsible for human cancers, the function of 4E-BPs in the inhibition of eukaryotic translation initiation,^{11,17–19} the significance of GW182 protein in the recruitment of the multiprotein machinery necessary for microRNA-mediated gene silencing,^{20–22} or the importance of Tau, FUS, and α -synuclein proteins in neurodegenerative diseases.^{23,24} Because the elastic properties of these biomolecules are responsible for the proper functioning of IDPs in the cellular context, i.e., for the association of complexes and the formation of biomolecular

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condensates via liquid–liquid phase separation such as, e.g., RNA-processing membraneless organelles,^{25,26} much attention has been paid to the hydrodynamic properties of IDPs. Experimental techniques, such as analytical ultracentrifugation (AUC), size exclusion chromatography (SEC), pulsed-field gradient nuclear magnetic resonance (PFG-NMR), dynamic light scattering (DLS), and fluorescence correlation spectroscopy (FCS), offer insights into hydrodynamic parameters (as reviewed by Bialobrzeski et al.²⁷). However, due to the distinct limitations of each experimental approach, ongoing research aims to devise phenomenological methods for calculating the hydrodynamic radius (R_h). These methods may involve deriving R_h from the radius of gyration (R_g) determined by small-angle X-ray scattering (SAXS)^{28,29} or exploiting the conformational backbone propensity of IDPs.^{30,31} However, it has recently been noted that inferring structural properties of the IDP conformational ensembles from SAXS is prone to a high degree of uncertainty.³²

A theoretical Monte Carlo approach was also developed on the basis of a bead chain model showing that proper consideration of the excluded volume effect is critical for estimating the R_h value of the disordered N-terminal Sic1 fragment,³³ in accordance with FCS experimental results.³⁴

Simultaneously, significant effort is being invested in developing numerical models that extract the characteristics of IDPs from conformational ensembles obtained using molecular dynamics (MD) simulations, deep learning, or energy minimization algorithms.^{35–47} However, the molecular flexibility of IDPs introduces substantial complexities when determining their hydrodynamic properties. Two main issues here are the large number of degrees of freedom and the long time scales of relaxation of the internal coordinates of the molecules. These factors prohibit direct calculation of the experimentally relevant long-time diffusion coefficient from either molecular or Brownian dynamics trajectories. One popular approximation that circumvents this difficulty is to assume that the macromolecule is rigidly frozen in one of a large number of possible conformations. Transport properties are then calculated by treating the molecule as a rigid body, and the results are averaged over an equilibrium ensemble.^{48–51} Nevertheless, the validity and accuracy of this approximation remain uncertain. Additionally, the generation of conformational ensembles can be a bottleneck for long chains (beyond ~300 amino acid residues) because it requires time-consuming MD simulations and/or the construction of new databases of short peptide conformations.

There is, therefore, a strong need to develop a numerically efficient solution that would enable reliable calculation of the long-time diffusion coefficient of any long chain IDP, such as one with 1000 amino acid residues, solely on the basis of its sequence information.

In this study, we introduce a new theoretical approach for both generating conformational ensembles of IDPs and calculating their hydrodynamic properties. This method enables a swift estimation of the diffusion coefficient for long IDPs in a matter of minutes, with superior accuracy compared to that of existing methods. This assertion is substantiated through rigorous testing of the model on a diverse set of experimental results obtained for 43 proteins. The data set includes both literature data and R_h values measured for a set of new IDP constructs using FCS under mild conditions (see the [Supporting Information](#)).

We present our results in terms of the hydrodynamic radius of a molecule, R_h . This radius represents the size of a solid sphere that possesses the same translational diffusion coefficient, D , as the given molecule under identical buffer conditions. Therefore, $R_h = k_B T / 6\pi\eta D$, where T is the temperature and η is the viscosity.

An important observation by Fixman^{52,53} is that the diffusion coefficient of a flexible macromolecule is time-dependent, with well-defined short- and long-time limits. The disparity between the two is attributed to the effects associated with relaxation of the internal coordinates of the molecule, as well as rotation of the macromolecule as a whole.^{52,54,55} The positivity of the dissipation rate in the system implies that the long-time diffusion coefficient (D_l) is always smaller than the short-time diffusivity (D_s).⁵³ The focus of theoretical approaches should be the determination of the former quantity, as it is the one measured in experiments utilizing techniques like FCS, AUC, or DLS. Unfortunately, the calculation of D_l is significantly more challenging than that of D_s because it involves the computation of time-dependent quantities, such as the memory function, which describes the relaxation effects. An additional point to keep in mind is that the value of the short-time diffusion coefficient depends on the choice of the point that one tracks.^{55–58} In contrast, the long-time diffusivity is independent of the choice of reference point.⁵⁹

The methods for predicting the diffusion coefficient can be broadly split into three categories: atomistic, phenomenological, and coarse-grained. For small proteins, high-resolution, atomistic MD methods can be used,⁶⁰ but they require either simulating the surrounding water molecules explicitly, which is very computationally intensive, or an implicit solvent scheme. In the case of implicit solvent methods, addressing hydrodynamic interactions between distant parts of the molecule^{61–64} and thermalization⁶⁵ pose significant challenges. Additionally, even for the smallest proteins, it is prohibitively difficult to obtain statistically meaningful data over the 10–100 ms scale, which would enable the direct computation of the long-time diffusion coefficient.

The other extreme consists of phenomenological models that predict R_h from the number of residues N and possibly other parameters, such as the total charge or amino acid composition. Theoretical considerations of Rouse, who modeled a protein as a Gaussian chain,⁶⁶ provided a foundation to the power law relationship $R_h \sim N^{1/2}$. The classical Rouse model employs random displacements between the monomers. If we assume complete independence of displacements between each consecutive pair of monomers, the central limit theorem dictates that as N approaches infinity, the squared end-to-end distance should conform to a scaled $\chi^2(3)$ distribution. Consequently, the dimensions of such an idealized chain are expected to scale with \sqrt{N} . Later work of Zimm included the effect of excluded volume,⁶⁷ which resulted in the scaling $R_h \sim N^\gamma$ with $\gamma = 0.588$.

Phenomenological size–length relationships that include other variables involve a number of fitting parameters. As a result, their range of applicability outside of the fitting data set is difficult to assess. An alternative phenomenological approach proposed by Pesce et al.²⁹ employs the radius of gyration obtained from SAXS experiments to estimate R_h . This is substantiated by the observation that within the Kirkwood–Riseman approximation⁶⁸ R_h and R_g share the same scaling relationship with N as long as the pair-displacement distribution converges under appropriate scaling to a Gaussian for large N values.

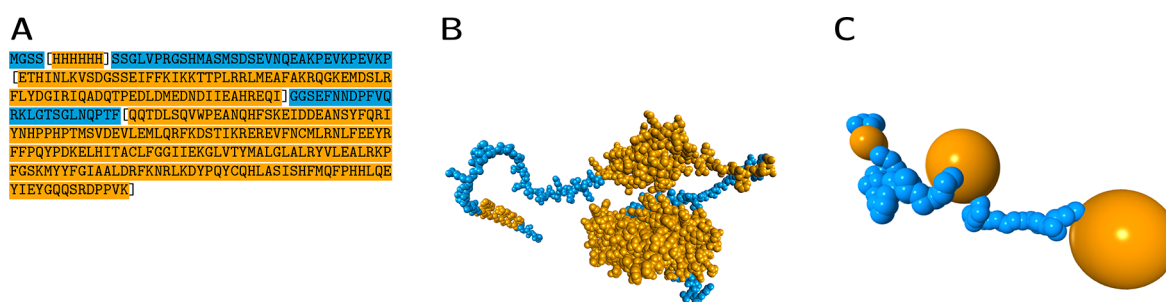


Figure 1. Construction of the coarse-grained globule-linker model (GLM) for an illustrative IDP, H₆–SUMO–CNOT1 (800–999), containing three ordered domains of different sizes (no. 28 in Table S1). (A) Sequence with highlighted ordered (orange) and disordered (blue) segments, and domain boundaries marked by square brackets. (B) Representative full atom conformation generated by AlphaFold2 (for visualization purposes only;^{69,70} beads with van der Waals radii; hydrogen atoms omitted for the sake of clarity). Ordered clusters (orange) form dense blobs connected with linkers (blue). (C) Visualization of a representative configuration generated using the GLM method in which beads are displayed with their hydrodynamic radii.

Finally, coarse-grained models, like our method, employ larger units (typically one or two per amino acid residue) as building blocks for the structure prediction scheme, along with approximate interaction potentials between subunits, to simulate the equilibrium ensemble of configurations for a given molecule. These configurations are then combined with an approximation of the hydrodynamic properties to compute the diffusion coefficient. Essentially, the computation of the latter for elastic macromolecules addresses two interconnected challenges: predicting the conformations of molecules on the basis of available biochemical data and then using these conformations to predict hydrodynamic properties.

The different exponents in the power law relationships of Rouse⁶⁶ and Zimm⁶⁷ demonstrate that even the most basic method for approximating configurations must take into account excluded volume interactions.

A software that can accommodate excluded volume interactions for a disordered chain is Flexible Meccano (FM).³⁷ In addition to volume exclusion, it considers the distribution of Ramachandran angles determined from crystallographic protein structures when sampling conformations. However, FM treats the entire chain as unstructured, so it cannot be used to model proteins that possess both globular and unfolded segments, which are in fact much more common than fully unstructured chains. Unfortunately, FM has a closed license that precludes necessary modifications to accommodate folded regions of proteins.

The complex angle distributions used by FM are crucial when computing NMR parameters that are sensitive to short-range details of the pair distribution function, such as residual dipolar couplings, paramagnetic relaxation enhancement, or *J* coupling. However, upon closer examination, the pair-distance distribution generated by FM and a simpler model presented in this paper, globule-linker model (GLM; described below), become virtually identical for amino acids separated by >15 residues along the chain.

The highly localized differences between structures at small sequential distances have a minimal influence on the estimations of R_h . It is important to recall that for amino acid residues separated by a distance r , the dipolar coupling decays as r^{-3} , while the decay rate of hydrodynamic interactions (HI) is only r^{-1} . Therefore, HI are long-range and less sensitive to near-neighbor distributions, with contributions to the diffusion coefficient of near neighbors and far neighbors being $O(N)$ and $O(N^{2-\gamma}) = O(N^{1.4})$, respectively.

Guided by these considerations, we have implemented the simplest extension of Zimm's chain, the globule-linker model (GLM), designed to comprehensively represent IDPs that contain globular domains connected by unstructured fragments. In particular, the GLM approach reflects the idea that the hydrodynamic radius corresponding to the experimentally measured long-time diffusion coefficient can be predicted under a minimal model that incorporates knowledge of domain boundaries in long protein chains and excluded volume interactions. In the model (Figure 1A–C), we represented the protein as an assembly of spheres of different sizes. Within the GLM approach, the conformational sampling is split into four stages: selection of domain boundaries, computation of steric radii of approximating spheres for globular domains, generation of locations of the domains and linkers, and addition of the hydration layer to the linkers.

First, the protein sequence fragments to be treated as folded domains and mimicked by larger beads within GLM are selected using disorder probability P predicted by Disopred3.⁷¹ A fragment is assumed to be ordered if the P value is <50% for at least three subsequent amino acid residues, and the ordered fragments within a single folded domain can be linked by loops, whose length does not exceed 14 residues.⁷² Because Disopred3 has been trained on the experimental data sets to obtain position-specific scores calculated for each amino acid residue,⁷¹ the P value involves implicitly the sequence specificity, reflecting the intramolecular interactions responsible for domain folding. Together with taking into account the experimentally established limit for the loop length,⁷² this approach enables us to create a biochemically relevant semiempirical model of globular domain boundaries. Such a globule boundary-annotated amino acid sequence is passed to the next stage of the modeling pipeline.

Second, the steric sizes of the approximating beads are computed. The structured domains are represented by a single larger sphere each, with the size depending on their mass m computed with the equation $R_h = (3m/4\pi\rho_{\text{globular}})^{1/3} + a_{\text{hydration}}$, where $\rho_{\text{globular}} = 0.52 \text{ Da}/\text{\AA}^3$,⁷³ with a single layer hydration shell taken to be $a_{\text{hydration}} = 3 \text{ \AA}$ thick. In the case of unstructured linkers between the domains, the beads representing amino acid residues of the linker are presumed to be indistinguishable. The composition of such linker sequences is known to be statistically biased toward the disorder-promoting residues (Pro, hydrophilic and charged residues) and deficient in hydrophobic and aromatic residues.^{74,75} The significance of the composition–conformation relationship was analyzed for IDPs in great detail

in terms of polar, polyampholytic, and polyelectrolytic tracts with different charge patterning (reviewed by Das et al.⁷⁵). Although it is clear that the dimensions of the charged IDP as a whole can be significantly influenced by electrostatic interactions depending on the solution conditions⁷ or charge patterning,⁷⁶ it seems reasonable to assume that, in solutions providing both sufficient hydration and ionic strength, the interactions between the polar and charged residues within the unstructured linker become less pronounced due to effective screening, and the exact pairwise potentials between the linker residues can be neglected. Each unstructured segment of length N is thus modeled as a chain of N identical spheres, each with a diameter equal to the C_α – C_α distance, and we obtain a list of steric radii of beads, which is passed on to the next modeling step.

Third, the centers of the beads are randomly sampled according to a generalization of a self-avoiding random walk. The distribution can be defined by first considering an auxiliary distribution of random walks of chains of spheres defined by demanding that distances between the centers of consecutive spheres along the chains are equal to the sum of their respective radii, and that each vector joining centers of adjacent spheres has a spherically uniform distribution. We then define the self-avoiding random walk of spheres (SARWS) to have the sphere centers distributed according to the random walk of spheres, conditional on the absence of self-intersections. Sampling from this distribution is achieved by a recursive algorithm described in the [Supporting Information](#), which offers accelerated sampling as compared to a one-by-one randomization. The SARWS algorithm ensures that the excluded volume of the chain is accounted for.

The fourth and final stage of the conformer generation process takes in the locations of the centers of the spheres generated in the previous step and adjusts their size to better reflect the hydrodynamic thickness of the linkers. We transformed the sampled conformations into a hydrodynamic model by increasing bead sizes in the disordered fragments of generated conformations to an $R_{\text{disordered}}$ of 4.2 Å, corresponding to the median value for all amino acids.⁷⁷ In the resulting hydrodynamic model of linkers, the neighboring beads show substantial overlaps, requiring a careful treatment of the mobility matrices (see ref 78 for details). Note that the value of $R_{\text{disordered}}$ has an only minor impact on the final results, because the hydrodynamic radius of long slender filaments depends logarithmically on their thickness.^{79–82}

To compute R_h from the estimated ensembles, we have implemented two algorithms: the Kirkwood formula and the minimum dissipation approximation (MDA) method of Cichocki et al.⁵⁹ Within the first approach,⁸³ the hydrodynamic radius of a macromolecule is approximated by

$$\frac{1}{R_h^{\text{K}}} = \frac{1}{N^2} \sum_{i=1}^N \left(\frac{1}{a_i} + \sum_{j=1, j \neq i}^N \left\langle \frac{1}{r_{ij}} \right\rangle \right) \quad (1)$$

where N is the total number of beads in the IDP model, a_i is the hydrodynamic radius of bead i , $r_{ij} = |\mathbf{r}_j - \mathbf{r}_i|$ is the distance between beads i and j , and the angle brackets denote the average over the equilibrium ensemble. One can show that this corresponds to the ensemble-averaged short-time diffusion coefficient of the geometric center of the macromolecule, $\mathbf{r}_c = N^{-1} \sum_{i=1}^N \mathbf{r}_i$. Note that the geometric center fluctuates as the shape of the molecule evolves and does not correspond to any

fixed position within it. A simplified form of the Kirkwood formula is often used^{42,84,85}

$$\frac{1}{R_h^{\text{K}}} \approx \frac{1}{N^2} \sum_{i=1}^N \sum_{j=1, j \neq i}^N \left\langle \frac{1}{r_{ij}} \right\rangle \quad (2)$$

where the single-bead terms $1/a_i$ are dropped, as their contribution becomes negligible in the large N limit. This is the form that we will use in the work presented here.

A better estimate of R_h , corresponding to the long-time diffusion coefficient, requires a more in-depth description of the hydrodynamic interactions between the beads. To this end, one introduces mobility matrix $\boldsymbol{\mu}$,⁵⁴ which links the velocities of the beads with the forces acting on them, according to

$$\mathbf{U}_i = \sum_j \boldsymbol{\mu}_{ij} \mathbf{F}_j \quad (3)$$

where \mathbf{U}_i is the velocity of bead i , whereas \mathbf{F}_j is the force with which bead j acts on the fluid. On the basis of the mobility matrix, one defines a matrix \mathbf{A} indexed by the bead labels (i, j) , $A_{ij} = 2\pi\eta \text{Tr}(\boldsymbol{\mu}_{ij})$ and its inverse $\mathbf{B} = \mathbf{A}^{-1}$. One can then construct the MDA⁵⁹ for R_h as

$$R_h^{\text{MDA}} = \sum_{ij} B_{ij} \quad (4)$$

Note that eq 4 is general and can be used for different models of hydrodynamic interactions, both simple models (e.g., Rotne–Prager far-field approximation⁸⁶) and more sophisticated approaches, like the multipole expansion method.^{87,88} In this work, we use the generalized Rotne–Prager approximation to calculate the mobility matrix, as described in refs 89–91. This approximation is now also available as a Python package, `pygrpy`.⁹² For non-overlapping beads, the elements of matrix \mathbf{A} have then a particularly simple form: $A_{ij} = \langle 1/r_{ij} \rangle$ for $i \neq j$, and $A_{ii} = 1/a_i$. The formulas for overlapping beads can be found in the [Supporting Information](#).

The MDA corresponds to the calculation of the short-time diffusion coefficient of the diffusion center of a molecule,⁵⁸ which is a point inside the molecule where D_s is minimal. The position of the diffusion center is $\mathbf{r}_d = \sum_{i=1}^N x_i \mathbf{r}_i$, with the weights given by $x_i = \sum_j B_{ij} / \sum_{k,j} B_{kj}$. Because D_s is always larger than its long-time counterpart, D_l , MDA provides the best estimation for the long-time diffusion coefficient of all of the methods that utilize D_s for this purpose. The MDA turns out to be more robust when dealing with large differences in the sizes of beads used to model constituent parts of the macromolecule, because in such cases the equal weights of the geometric center of the macromolecule used in the Kirkwood formula differ significantly from the optimal weights of the diffusion center.

We combined each method of generating conformers with each method of computing R_h , which resulted in four different theoretical approaches, the predictions of which (Table S2) were then compared with experimental data. For this purpose, we have obtained 15 new IDP constructs covering a wide range of chain lengths, folded domain contents, and charge states and determined their R_h using FCS (Figure 2 and Figures S2–S6; for further experimental details, see the [Supporting Information](#)).

The experimental benchmark set (Table S1) was thus composed of both the new FCS measurements and R_h values selected from the literature on the basis of the following criteria. The proteins had sequences that could be unambiguously identified in the literature or in the UniProtKB database and

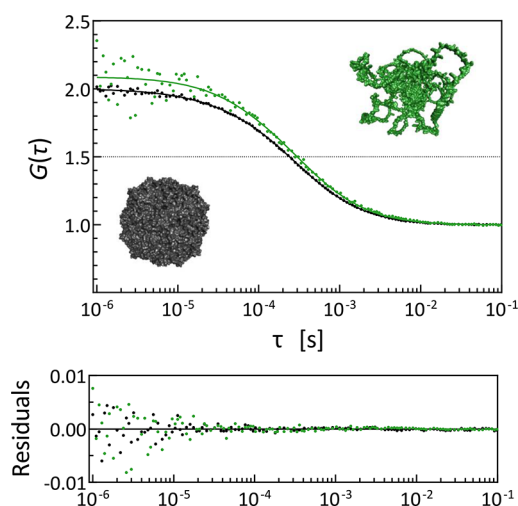


Figure 2. Examples of normalized FCS autocorrelation curves with raw fitting residuals for an intrinsically disordered H_6 -SUMO-GW182SD-mCherry ($N = 809$; $R_h = 66 \pm 6$ Å) (green) in comparison with apoferritin ($N = 4200$; $R_h = 58 \pm 3$ Å) (black). The crystal structure of apoferritin (Protein Data Bank entry 2w0o⁹³) and the putative conformation of H_6 -SUMO-GW182SD-mCherry predicted by AlphaFold⁶⁹ are shown for the purpose of illustration, preserving the relative sizes of the solvent accessible surfaces of atoms.

were measured under well-defined, mild conditions (temperature of 20–26 °C, buffer of pH 7–8, and ionic strength corresponding to 75–300 mM NaCl), and their hydrodynamic radii were determined directly from appropriate experiments without conversions from other experimental quantities, such as R_g .^{29,94–111} This is, to our best knowledge, the largest benchmark set encompassing experimental R_h values for 38 IDPs and six globular model proteins, measured under comparable conditions (Figure 3).

The results of tests performed for our four theoretical approaches against the benchmark set are listed in Table 1, and Figure 4 shows a visual comparison of the deviations between theory and experiment. Additionally, we provide various power law fits^{112–114} for comparison of the prediction accuracy (Table S3).

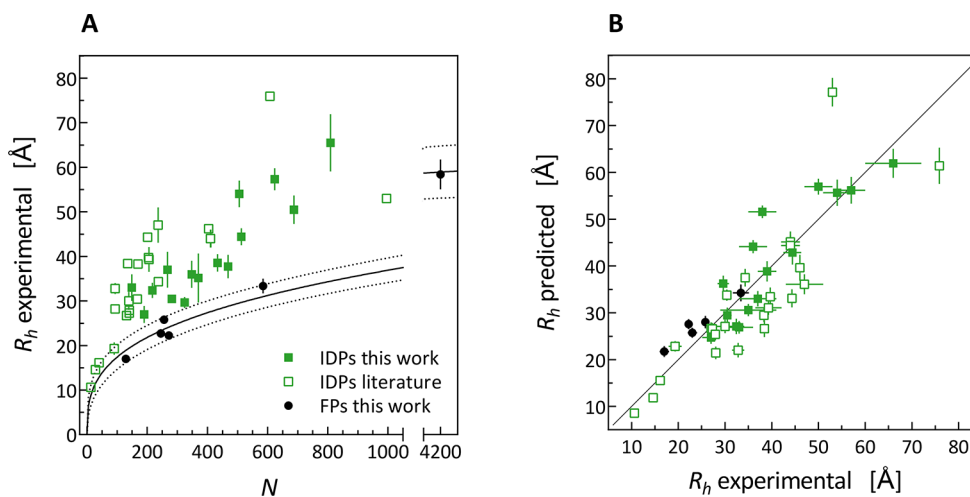


Figure 3. (A) Experimental R_h values plotted vs the number of amino acid residues in the protein chain, N , and power law curve fitted to R_h values of folded proteins (FPs) together with the 95% confidence band. (B) Direct comparison of the predicted vs measured R_h values for all of the proteins modeled using the MDA+GLM approach.

We compare the accuracy of the previous and new model under six metrics (Table 1): the square root of the mean square deviation (RMSD), the square root of the mean square relative deviation (RMSRD), Pearson's coefficient (R^2), Pearson's coefficient adjusted for fitting parameters (R_{adj}^2), the third quartile of the absolute error (Q_3^{AE}), and the third quartile of the relative error (Q_3^{RE}). Whenever a fitting procedure is required, we use leave-one-out cross-validation to compute error metrics. We also have chosen to test the relative deviations to reduce the undue weight given to the new, very long sequences in our data set. Similarly, outlier-robust metrics of the third quartile were included to reduce the impact of a single-sequence misprediction on the final comparisons. In all evaluation metrics, the MDA+GLM approach performs the best. Surprisingly, it is the only model that performs better than the power law baseline in any of the evaluation metrics.

Interestingly, it is apparent from the comparison of the results obtained using MDA+GLM with those from MDA+GLM(ND) in Table 1 and Figure 4 (A and B) that the proper identification of the globular domain boundaries proves to be the main condition for successfully estimating the R_h value of an IDP, with better accuracy than all other tested approaches. This means that the pairwise interactions between the linker amino acid residues influence R_h to a lesser extent, while the sizes of the globular domains and their relative spatial distribution are very important.

It should be mentioned, however, that a significant contribution to the discrepancies between the experimental and predicted R_h values (Figure S8) comes from the intrinsic properties of the individual experimental methods, which suffer from typical errors or limitations and are usually not taken into account when reporting the final experimental results. PGF-NMR measurements are the most unambiguous and accurate, but their effective application is limited to smaller proteins (up to 200–300 amino acid residues long) at high concentrations. It is worth noting that the agreement of the values of R_h predicted by MDA+GLM with the PGF-NMR results is excellent (Figure S8C). FCS is the only method that addresses the self-diffusion of molecules at the low-concentration limit. Raw FCS measurements can be refined to exclude possible oligomerization or aggregation during the experiment on the basis of the count

Table 1. Comparison of Error Statistics of Various Models^a

model	n_{fp}	RMSD (Å)	RMSRD (%)	R^2	R^2_{adj}	Q_3^{AE} (Å)	Q_3^{RE} (%)
MDA+GLM	0	7.09	18.15	0.71	0.71	6.80	22.51
MDA+GLM(ND)	0	9.48	28.02	0.48	0.48	11.88	29.31
KR+GLM	0	12.82	34.69	0.05	0.05	17.59	42.95
KR+GLM(ND)	0	9.25	27.31	0.50	0.50	11.11	29.44
random coil	1	9.60	27.71	0.47	0.45	10.28	33.69
power law	2	8.46	24.80	0.59	0.56	9.63	26.08
power law (ref 112)	2	12.01	36.94	0.16	0.12	14.37	39.51
PPII-based (ref 30)	3	17.25	49.09	-0.72	-0.86	20.62	59.54
PPII and IQL-based (ref 31)	5	18.97	47.71	-1.08	-1.36	19.60	54.77
sequence-based (ref 112)	7	22.90	50.78	-2.05	-2.66	19.59	58.32

^a n_{fp} is the number of fitting parameters. ND indicates no domain information. Q_3 is the third quartile.

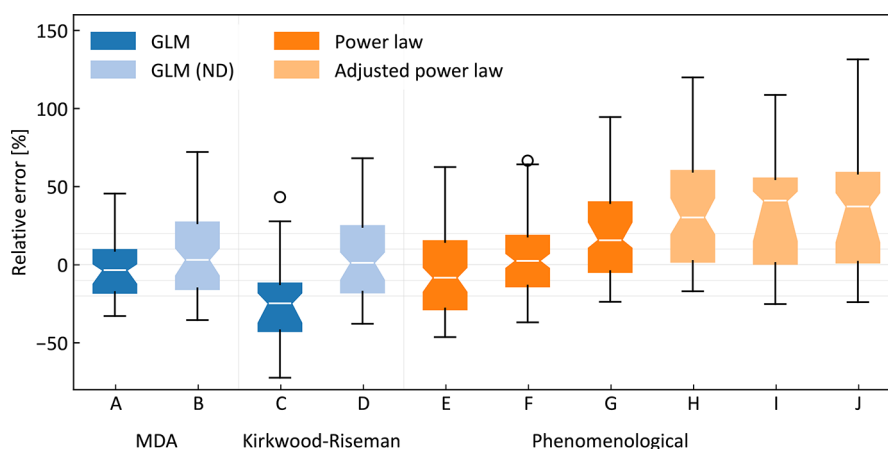


Figure 4. Comparison of different methods of estimation of R_h . Boxes show interquartile ranges with median confidence bands marked by notches. MDA with GLM ensemble generation (A) performs best on the IDP benchmark set with standard errors of 18.15% and 7.09 Å (compared to 24.80% and 8.46 Å for a simple power law). Methods based on the Kirkwood–Riseman R_h estimation (C and D) typically underestimate hydrodynamic size of the molecule. Power law fits with one free parameter (E) and two free parameters (F) evaluated using leave-one-out cross-validation are compared with the formerly reported power law¹¹² (G) and models based on polyproline II structure propensities without³⁰ (H) and with³¹ (I) regard to the charge, and a sequence-based model¹¹² (J) that takes into account the total charge of the molecule. Theoretical methods with no knowledge of the presence of domains in the IDP (ND; B and D) significantly overestimate the hydrodynamic size of the molecule. Domain data can be incorporated into our ensemble generation engine leading to more accurate estimates of R_h (A). Note that experimental uncertainty also contributes to the errors presented here and in Table 1.

rates, but it is impossible to avoid proteolytic instability of proteins and, consequently, the appearance of impurities with a lower molar mass, which may potentially result in apparently lower values of R_h (Figure S8B). On the contrary, SEC is the easiest approach for removing lower-mass impurities, but it involves diffusion of molecules at higher concentrations through a medium with pores of a specific shape under the influence of pressure. An additional common disadvantage is calibration based on R_h of standard proteins determined under various conditions and the lack of appropriate propagation of the calibration experimental uncertainty. Consequently, SEC measurements can be highly scattered (Figure S8D). The largest outlier in our analysis concerns R_h determined using SEC for fesselin without providing experimental uncertainty (Id. 43, Tables S1 and S2 and Figures S7 and S8D). The DLS method is the most prone to overestimating experimental values (Figure S8E), because the presence of even a small number of aggregates with a larger molar mass generates a huge contribution to the intensity of scattered light. Finally, AUC yields sedimentation coefficients, and their interpretation in terms of exact values of R_h requires some assumptions that are not obvious for IDPs, such as, e.g., partial specific protein volume.¹¹⁵ The second largest outlier in our set is the OMM-64 protein (Id. 39, Tables

S1 and S2 and Figures S7 and S8F) with the R_h value determined using AUC, which is very close to the power law curve for completely denatured proteins.¹¹⁶

In conclusion, we have presented a simple, first-principles model for the prediction of R_h without any fitting parameters and achieved favorable comparison with a large benchmark set. The sizes and positions of the globular domains proved to be the dominating factors that influence the hydrodynamic properties of the IDP chain as a whole. Moreover, due to the relative simplicity of the model, all of the calculations for a given protein can be performed in ~ 1 min on a typical laptop, which is contrasted with MD simulation-based conformer generation methods that require supercomputers and take many days. Moreover, the MDA+GLM approach demonstrates satisfactory convergence even with ensemble sizes as small as 40 conformers (Figure S1).

Our benchmark set, in which the previously known IDPs were complemented by a set of newly obtained proteins, constitutes a significant step forward in predicting the hydrodynamic properties of IDPs. It includes a higher degree of conformational variety, with a stronger emphasis on multidomain proteins, longer chains, and a much wider range of charge states compared to the reference sets used previously.^{30,112} This diversity allows

for more reliable testing of theoretical models. In particular, the presence of large polyanionic proteins in our set revealed that the R_h values obtained using phenomenological models corrected to account for the absolute net charge seem to be overestimated [Figure 4 (I and J)].

The sequence specificity effects are neglected in our model for the linker fragments, which is one of the possible sources of uncertainty. However, in our opinion, it is an acceptable level of error for such a quick numerical method. Further developments of the MDA+GLM approach are needed to take into account the dependence of R_h on the environmental conditions^{6–8} and the formation of complexes. More subtle effects related to the conformational properties of the linkers can be also included using sequence-based conformational ensembles.^{45,47,117} Nevertheless, our results demonstrate that the relatively simple globule-linker model for conformational ensemble construction, in combination with the minimum dissipation approximation, can serve as the starting point for developing further phenomenological corrections. These improvements could incorporate factors such as amino acid sequence composition, residue charge, and counterion binding. When using the MDA+GLM approach, all excluded volume effects are already correctly accounted for, with any further deviations hinting at the interesting physical and chemical properties of the molecules.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acs.jpcllett.4c00312>.

Experimental R_h values, Figures S1–S8, and additional methodological details (PDF)

Theoretical R_h values from GLM, phenomenological R_h values, and protein sequences with marked domains (Tables S2–S4) (XLSX)

Transparent Peer Review report available (PDF)

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Notes

The authors declare no competing financial interest.

The programs referenced in this article can be found on GitHub. For the sake of convenience, an API and a command-line Python utility, `glm_mda_diffusion`, have been provided.¹¹⁸ This package builds upon previous ones, namely, `pychastic` and `sarw_spheres`, both of which are also accessible on GitHub.

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