Integrating multiple experimental data to determine conformational ensembles of an intrinsically disordered protein

Gregory-Neal. W. Gomes¹,²*, Mickaël Krzeminski³,⁴, Erik. W. Martin⁵, Tanja Mittag⁵, Teresa Head-Gordon⁶, Julie D. Forman-Kay³,⁴, and Claudiu C. Gradinaru¹,²*

¹Department of Physics, University of Toronto, Toronto, Ontario M5G 1X8, Canada
²Department of Chemical and Physical Sciences, University of Toronto Mississauga, Mississauga, Ontario L5L 1C6, Canada
³Molecular Medicine Program, Hospital for Sick Children, Toronto, Ontario M5S 1A8
⁴Department of Biochemistry, University of Toronto, Toronto, Ontario M5G 1X8, Canada
⁵Department of Structural Biology, St. Jude Children’s Research Hospital, Memphis, TN 38105, USA
⁶Departments of Chemistry, Bioengineering, Chemical and Biomolecular Engineering University of California, Berkeley, California 94720, United States
*Authors to whom correspondence should be addressed
Abstract

Intrinsically disordered proteins (IDPs) have fluctuating heterogeneous conformations, which makes structural characterization challenging. Although challenging, characterizing the conformational ensembles of IDPs is of great interest, since their conformational ensembles are the link between their sequences and functions. An accurate description of IDP conformational ensembles depends crucially on the amount and quality of the experimental data used in their calculations. We used integrative modelling to understand and implement conformational restraints imposed by the most common structural techniques for IDPs: Nuclear Magnetic Resonance (NMR) spectroscopy, Small-angle X-ray Scattering (SAXS), and single-molecule Förster Resonance Energy Transfer (smFRET). Agreement with such a diverse set of experimental data suggests that details of the generated ensembles can be examined with a high degree of confidence. Using the disordered N-terminal region of the Sic1 protein as a test case, we examined relationships between average global polymeric descriptions and higher-moments of their distributions. To resolve apparent discrepancies between smFRET and SAXS inferences, we integrated SAXS data with non-smFRET (NMR) data and reserved the smFRET data as an independent validation. Consistency with smFRET, which was not guaranteed a priori, indicates that, globally, the perturbative effects of NMR or smFRET labels on the Sic1 ensemble are minimal. Analysis of the integrative ensembles revealed distinguishing features of Sic1, such as overall compactness and large end-to-end distance fluctuations, which are consistent with biophysical models of Sic1’s ultrasensitive binding to its partner Cdc4. Our results underscore the importance of integrative modelling in calculating and drawing conclusions from IDP conformational ensembles.
1 Introduction

Under physiological conditions, the amino acid sequences of intrinsically disordered proteins (IDPs) encode for a large and heterogeneous ensemble of conformations, allowing them to perform critical biological functions[1, 2]. The properties of IDP conformational ensembles are intimately related to their function in health and disease[3]. This has prompted intense efforts to develop formal and heuristic descriptions of how sequence properties relate to conformational ensembles[4–8], and how the properties of conformational ensembles, once determined, can be mined to generate hypotheses about biological function[9–11]. Conformational ensembles are therefore central to understanding both sequence-to-ensemble and ensemble-to-function relationships in IDPs, which makes their accurate and comprehensive characterization of high importance.

To provide insights into the structural properties of IDPs, Nuclear Magnetic Resonance (NMR)[12], Small-Angle X-Ray Scattering (SAXS)[13], and single-molecule Förster Resonance Energy Transfer (smFRET)[14, 15] have emerged as particularly powerful techniques. Computational approaches to integrate the information from these measurements typically represent conformational ensembles as a collection of structures, each described by its atomic coordinates, and use the experimental data for constructing (e.g., restraining or re-weighting), or validating the ensemble calculation[16–18].

Despite their demonstrated complementarity[19–21], conformational ensembles which use data from all three techniques in their construction or validation are rarely reported. Aznauryan et al., reported ensembles of ubiquitin denatured in 8 M urea which are consistent with SAXS and a large number of restraints from NMR and smFRET experiments[20]. However, concerns about the mutual consistency of smFRET and SAXS data posit that in the absence of denaturant, the FRET fluorophores could interact with each other and/or the IDP itself[22]. Piana et al., reported ensembles of α-synuclein in physiological conditions, which are directly compared to SAXS and NMR data, but are compared to distances inferred from smFRET data using an assumed homopolymer model[23]. However, it is difficult to determine which, if any, homopolymer model is appropriate for a particular heteropolymeric IDP[24, 25]. Thus, using data from all three techniques to construct or validate conformational ensembles of an IDP (i) in physiological conditions and (ii) without assuming a homopolymer model, would provide valuable insights into each technique’s sensitivity to different aspects of IDP structure.

Polymer physics offers a conceptual framework for describing the conformationally heterogeneous ensembles that are determined using the data from these experiments [14, 15, 26, 27]. Considerable attention has focused on the calculation/measurement of first moments of global polymeric descriptions, such as ensemble-averaged values for the radius of gyration \( R_g \), end-to-end distance \( R_{ee} \), and asphericity/shape. In particular, recent studies have reported on the determination of one or more of these values from single measurements[22, 24, 28–30]. It is becoming increasingly clear however, that for IDPs, which are bona fide heteropolymers, conclusions drawn from first-moments, particularly those drawn from single measurements, risk over-interpretation[21, 24, 28, 31, 32]. The accuracy of entire distributions of these polymeric parameters, and their joint distributions, determines the extent to which sophisticated sequence-to-ensemble relationships can be developed[32]. Achieving a high level of accuracy will require integrative modelling, and will depend crucially on the amount and quality of the experimental data used in ensemble calculations.

We therefore sought to determine conformational ensembles of an IDP in physiological conditions with conformational restraints/validation imposed by NMR, SAXS, and smFRET. Using the disordered N-terminal region of the Sic1 protein as a test case (see below), we generated new smFRET and SAXS data to complement previously published NMR data[33, 34]. To combine these datasets, we used the ENSEM-
BLE approach (Fig. 1), which selects a subset of conformations from a large starting pool of conformations to achieve agreement with experimental data[17, 35, 36]. Our final ensembles of Sic1 are consistent with a diverse set of experimental data suggesting that their properties can be examined with a high degree of confidence. This allowed us to examine relationships between average global polymeric descriptions of Sic1 and higher-moments of their distributions. Our results underscore the importance of integrative modelling in determining ensembles, and therefore as a prerequisite for understanding sequence-to-ensemble and ensemble-to-function relationships.

Figure 1: A schematic showing the ENSEMBLE approach for SAXS and smFRET data from an ensemble of structures $X = [X_1, ..., X_k, ..., X_{N_{\text{conf}}}]$. (A-B) The SAXS intensity curve of each conformation $i_k(q)$ is back-calculated from the atomic coordinates using CRYSOL[37]. (C) The linear average of the CRYSOL-calculated SAXS profiles of individual conformers (black) is compared with the experimental SAXS profile (yellow). (D-E) Per-conformer FRET efficiencies $e_k$ are calculated from the mean distance between dyes $\langle r_{DA} \rangle_k$ predicted by accessible volume simulations[38, 39]. (F) The ensemble-averaged transfer efficiency $\langle E \rangle_{\text{ens}} = \langle e_k \rangle$ (grey vertical line in E and F) is compared to the mean experimental transfer efficiency $\langle E \rangle_{\text{exp}}$ (yellow vertical line collinear with grey line in F).

Achieving our objective of determining Sic1 ensembles consistent with all three datasets allows us to provide additional insight into the so-called “smFRET and SAXS controversy”[40–42]. Previous studies have either (i) posited attractive fluorophore interactions in the absence of denaturant[22, 29], or (ii) have jointly restrained ensemble calculations using both the smFRET and SAXS data[19, 21, 31]. The latter approach is based on the recognition that for heteropolymers, deviations from homopolymer chain statistics can cause smFRET and SAXS to be sensitive to different aspects of IDP structure (e.g., $R_{ee}$ and $R_g$ respectively)[21, 28]. For a given IDP and set of labels, both explanations for discrepant inferences are a priori plausible and so additional experimental information is needed. Additional experimental information in approach (ii) is provided by self-consistent smFRET distance inferences with labels of varying physicochemical properties[19] or self-consistent SAXS measurements of samples with and without FRET labels[21]. Rather, we provide additional experimental information in the form of NMR restraints, and reserve the smFRET data as an independent validation. Consistency with the smFRET data indicates that globally perturbative effects of
NMR or smFRET labels on the Sic1 ensemble are minimal. Our test case IDP, is the N-terminal 90 residues of the full-length disordered protein Sic1. In yeast, the disordered protein Sic1 is eliminated via ubiquitination by the SCF\textsuperscript{Cdc4} ubiquitin ligase and subsequent degradation by the proteasome, allowing initiation of DNA replication\cite{43, 44}. Sic1 binding to Cdc4, the substrate recognition subunit of the ubiquitin ligase, generally requires phosphorylation of a minimum of any six of the nine Cdc4 phosphodegron (CPD) sites on (full length) Sic1. This effectively sets a high threshold for the level of active G1 CDK required to initiate transition to S-phase. This ultrasensitivity with respect to G1 CDK activity ensures a coordinated onset of DNA synthesis and genomic stability\cite{43}. The N-terminal 90 residues of Sic1 (henceforth Sic1) are sufficient for targeting to Cdc4 when highly phosphorylated (henceforth pSic1), making this region a valuable model for structural characterization\cite{45}. Neither phosphorylation, nor binding to Cdc4 leads to folding of Sic1\cite{33, 34}. As the binding properties of Sic1 and pSic1 are vastly different, accurate conformational ensembles of Sic1 and pSic1 are central to developing and validating biophysical models of their differential binding\cite{46–48}.

2 Results

2.1 Measurements of $R_{ee}$ and $R_g$ inferred individually from smFRET or SAXS provide discrepant descriptions of Sic1 and pSic1 conformational ensembles

Fig. 2 A-C shows smFRET data measured on the Sic1 FRET construct, which is based on Sic1(1-90) and hereafter called Sic1. This construct was labelled stochastically at its termini with the FRET donor Alexa Fluor 488 and acceptor Alexa Fluor 647 (Förster radius $R_0 = 52.2 \pm 1.1$ Å, SI Appendix 1.7). The histogram is fit to a Gaussian function to extract the mean transfer efficiency $\langle E \rangle_{exp}$, which reports on the end-to-end distance distribution $P(r_{ee})$ (see SI Appendix 1.10 for more details). Multisite phosphorylated Sic1 (pSic1) was generated via overnight incubation with Cyclin A/Cdk2 resulting in predominantly 6- and 7-fold phosphorylated Sic1, with a minor population of 5-fold phosphorylated Sic1 (determined by ESI mass spectrometry). Upon phosphorylation, $\langle E \rangle_{exp}$ decreases from 0.42 to 0.36 indicating chain expansion.

An estimate of the root-mean-squared end-to-end distance $R_{ee}$ can be made from $\langle E \rangle$ by assuming $P(r_{ee})$ is described by a homopolymer model (SI Appendix 1.10). However, the smFRET data itself ($\langle E \rangle_{exp}$) does not suggest which (if any) homopolymer model is appropriate for a certain IDP. There is considerable flexibility in the choice of homopolymer model and in how to rescale the root-mean-squared inter-dye distance $R_{D,A}$ to $R_{ee}$, resulting in a range of $R_{ee}$. The inferred $R_{ee}$ is 61-65 Å for Sic1 and 66-72 Å for pSic1, suggesting multisite phosphorylation results in an approximately 10% increase in $R_{ee}$ (SI Appendix, Table S2). The smFRET data set, examined alone, suggests that Sic1 is 7-13% more compact than a self-avoiding random coil (RC) ensemble generated with the statistical coil generator TraDES for Sic1\cite{49, 50}.

To infer the root-mean-squared radius of gyration $R_g$ from $R_{ee}$ requires an additional assumption about the polymeric nature of system under study, namely the ratio $G = R_{ee}^2/R_g^2$. It has recently been shown that finite-length heteropolymeric chains can take on values of $G$ that deviate from the values derived for infinitely long homopolymers in either the $\theta$-state (Gaussian chains, $G = 6$) or excluded-volume (EV)-limit (self-avoiding walks, $G \approx 6.25$)\cite{21, 28, 31}. Application of polymer-theoretic values of $G$ to the smFRET inferred $R_{ee}$ results in $R_g$ 24-27 Å for Sic1 and 26-29 Å for pSic1 (SI Appendix Table S3). These inferred $R_g$ values are systematically smaller than those inferred from the SAXS dataset (see below), or from integrated ensemble modelling (see below), similar to previously reported discrepancies between smFRET and SAXS\cite{19, 21, 29}.
Figure 2: (A-B) smFRET efficiency ($E$) histograms of Sic1 (A) and pSic1 (B) labelled with Alexa Fluor 488 and Alexa Fluor 647 at positions -1C and T90C in TE buffer pH 7.5 150 mM NaCl. (C) Example SAW homopolymer $P(r_{ee})$ distributions (left vertical scale) for Sic1 (black, $R_{ee} = 62.5$ Å) and pSic1 (red, $R_{ee} = 67.8$ Å). The shaded underlying region shows the FRET distance dependence function $E(r_{ee})$ (right vertical scale). (D) Dimensionless Kratky plots of Sic1 (black) and pSic1 (red), normalized by initial intensity $I_0$ and the $R_g$ estimated from the DATGNOM fit of the distance distribution function. (E) Guinier plots of Sic1 (black) and pSic1 (red). The solid circles are the data points selected for fitting ($q_{max}R_g < 1.1$) and the solid lines show the Guinier fits using these data points. (F) The normalized distance distribution function $P(r)$ estimated by DATGNOM for Sic1 (black) and pSic1 (red).

Fig. 2 D-F shows SAXS data for Sic1 and pSic1. $R_g$ was estimated to be approximately 30 Å for Sic1 and 32 Å for pSic1 using the Guinier approximation, and from the distance distribution function $P(r)$ obtained using the indirect Fourier transform of the regularized scattering curve (Fig. 2 E&F and SI Appendix 2.1). A model of chain statistics does not need to be specified, however, these methods are limited in describing IDPs and unfolded proteins[13, 19]. For example, the expanded and aspherical conformations of IDPs lead to a reduced range of scattering angles in which the Guinier approximation can be applied without systematic error[19]. The degree of underestimation of $R_g$ increases as the maximum scattering angle $q_{max}$ increases, while decreasing $q_{max}$ reduces the number of points restraining the Guinier fit, which increases the uncertainty in $R_g$[19] (see also, SI Appendix, Table S4).

One solution to these limitations is to model the protein chain explicitly by generating ensembles of conformations. This is epitomized by the Ensemble Optimization Method (EOM) [52] and ENSEMBLE [35]. Both approaches select a subset of conformations from an initial pool of conformations, such that the linear average of the CRYSOL-calculated SAXS profiles of individual conformers is in agreement with the full experimental SAXS profile (Fig. 1 A-C). However, the techniques differ in their generation of the initial pool of conformations and in the algorithm and cost-function used to minimize the disagreement with experiment (SI Appendix 2.2). Despite their differences, both ensemble-based approaches fit the SAXS data equally well, and resulted in nearly identical $R_g$ values, which are similar to the “model-free” estimates (SI Appendix, Table S5). As was seen from the smFRET data, multisite phosphorylation results in chain expansion; this is in agreement with the SAXS data that indicates an approximately 6% increase in $R_g$.

Similarly, Riback and coworkers have recently introduced a procedure for fitting SAXS data by pre-
generating ensembles of conformations with different properties (specifically, the strength and patterning of inter-residue attractions) and extracting dimensionless “molecular form factors” (MFFs)[22, 29]. The properties of interest are then inferred from the ensemble whose MFF best fits the data. Using the MFFs generated from homopolymer or heteropolymer simulations results in similar $R_g$ to the aforementioned methods (SI Appendix, Table S6). Thus, $R_g$ is strongly determined by the SAXS data, such that differences in the construction and refinement of models leads to minor differences in $R_g$.

Since conformations are explicitly represented in the EOM, ENSEMBLE, and MFF methods, the $R_{ee}$ (and hence $G$) of the determined ensembles can be calculated. Although the various ensembles fit the SAXS data equally well, they have distinct values of $R_{ee}$, i.e., from 71-81 Å for Sic1 and from 71-87 Å for pSic1 depending on the method used (SI Appendix, Tables S5&6). These ensembles thus have $G$ values from 5.1-7.7 and 4.6-7.9 for Sic1 and pSic1, respectively (SI Appendix, Tables S5&6). Unlike $R_g$, the SAXS data does not uniquely determine $R_{ee}$ and $G$, independent of modelling approach. Naturally, the accuracy of those aspects of the ensemble not strongly determined by the SAXS data will depend on the initial conformer generation and the optimization/selection algorithms. This suggests that integrating additional experimental data will improve structural inferences.

Similarly, for homopolymer-based smFRET inferences, modelling flexibility lead to a 4-6 Å range of inferred $R_{ee}$ for Sic1, while the accuracy of $\langle E\rangle_{exp}$ ($\pm 0.02$), roughly corresponds to an uncertainty in the inferred $R_{ee}$ of $\pm 2$ Å. Likewise, $G$ cannot be determined from the data itself, and must be assumed a priori. It would therefore be desirable to back calculate $\langle E\rangle_{ens}$ from a structural ensemble that is restrained by additional experimental data and to compare $\langle E\rangle_{ens}$ and $\langle E\rangle_{exp}$, directly.

### 2.2 Ensembles jointly restrained by SAXS and NMR data are consistent with measured FRET efficiencies

We hypothesized that jointly restraining ensembles with non-smFRET internal distance restraints and SAXS data could result in ensembles with back-calculated mean transfer efficiencies, $\langle E\rangle_{ens}$, in agreement with the experimental mean transfer efficiency $\langle E\rangle_{exp}$. In addition to independently validating the calculated ensemble, this would provide compelling evidence that the smFRET and SAXS data sets are mutually consistent.

To provide non-smFRET information for joint refinement with SAXS data we used previously published NMR data on Sic1[33, 34]. Briefly, the NMR data consist of $^{13}$C$_\alpha$ and $^{13}$C$_\beta$ chemical shifts (CSs) from Sic1 and Paramagnetic Relaxation Enhancement (PRE) data from six single-cysteine Sic1 mutants using a nitroxide spin label (MTSL) coupled to cysteine residues in positions -1, 21, 38, 64, 83, and 90. We used the ENSEMBLE approach to calculate ensembles that are in agreement with the NMR and SAXS data (see Materials and Methods and SI Appendix 3). We used fluorophore accessible volume (AV) simulations[38] to back-calculate the mean transfer efficiency $\langle E\rangle_{exp}$ from the sterically accessible space of the dye attached to each conformation via its flexible linker (see Materials and Methods and SI Appendix 3).

The agreement of the experimental and back-calculated NMR and SAXS data was quantified using the reduced $\chi^2$ metric to identifying statistically significant disagreement with experimental data ($\chi^2 >> 1$) (considering experimental and back-calculation errors, see Materials and Methods and SI Appendix 3). As a structureless null-hypothesis we also include a random coil (RC) ensemble generated by TraDES for Sic1. This RC ensemble is shown to be in very good agreement with excluded volume (EV) homopolymer statistics (see below). Table 1 summarizes the goodness of fit for Sic1 ensembles under various restraint combinations.

SI Appendix Fig. S3 shows typical examples of TraDES RC and SAXS-restrained fits to the experimental...
SAXS profiles.

Table 1: Goodness of fit for Sic1 $N_{\text{conf}} = 500$ ensembles $^a$

<table>
<thead>
<tr>
<th>Restraints</th>
<th>$\chi^2$ PRE</th>
<th>$\chi^2 13\alpha$ CS</th>
<th>$\chi^2 13\beta$ CS</th>
<th>$\chi^2$ SAXS</th>
<th>$\langle E \rangle_{\text{exp}} - \langle E \rangle_{\text{ens}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>TraDES RC (none)</td>
<td>1.51</td>
<td>0.479</td>
<td>0.466</td>
<td>1.85</td>
<td>0.12</td>
</tr>
<tr>
<td>SAXS</td>
<td>2.06</td>
<td>0.470</td>
<td>0.395</td>
<td>1.01</td>
<td>0.15</td>
</tr>
<tr>
<td>PRE</td>
<td>0.230</td>
<td>0.544</td>
<td>0.608</td>
<td>13.5</td>
<td>-0.22</td>
</tr>
<tr>
<td>SAXS+PRE</td>
<td>0.261</td>
<td>0.376</td>
<td>0.394</td>
<td>1.12</td>
<td>0.01</td>
</tr>
<tr>
<td>SAXS+PRE+CS+(E)-filter</td>
<td>0.231</td>
<td>0.317</td>
<td>0.231</td>
<td>1.12</td>
<td>0.01</td>
</tr>
</tbody>
</table>

$^a$ Fit quality of $N_{\text{conf}} = 500$ ensembles derived by combining conformations from five independently calculated $N_{\text{conf}} = 100$ ensembles. Differences $|\langle E \rangle_{\text{exp}} - \langle E \rangle_{\text{ens}}| \leq \sqrt{\sigma_{E,\text{exp}}^2 + \sigma_{E,\text{ens}}^2} \approx 0.02$ indicate no disagreement between back-calculated and experimental mean transfer efficiencies (see Materials and Methods).

For the TraDES RC ensemble, there is no statistically significant disagreement with the CS data ($\chi^2 < 1$). However, the agreement with the PRE, smFRET and SAXS data is poor. Internal distances between specific residues are generally larger in the RC ensemble than are expected from the PRE and smFRET data. On the other hand, the radius of gyration of this ensemble ($R_g \approx 28 \, \text{Å}$) is slightly smaller than SAXS-only estimates ($R_g \approx 30 \, \text{Å}$). Although specific internal distances in this ensemble are not reproduced, the mean-squared sum over all internal distances in this ensemble is only slightly less than indicated by the SAXS data, as $R_g = \sqrt{\frac{1}{2n^2} \sum_{ij} (r_{ij}^2)}$.

When only the SAXS data are used as a restraint, the ensemble reproduces the SAXS curve very well. However, relative to the RC ensemble, the overall larger inter-residue distances in the SAXS-only ensemble further deteriorate the agreement with data reporting on specific inter-residue distances from PRE and smFRET.

When only the PRE data are used as a restraint, the agreement with the PRE data is achieved at the expense of not agreeing with all other observables. This ensemble reproduces specific inter-residue distances encoded by the PRE data, but not the overall distribution of inter-residue distances encoded by the SAXS data. A corollary of the $r^{-6}$ PRE weighting is that the PRE ensemble average is dominated by contributions from compact conformations[53]. Consistent with this, the PRE-only ensemble is much more compact ($R_g \approx 22 \, \text{Å}$) than expected from the SAXS data. Similarly, the transfer efficiency calculated from the ensemble $\langle E \rangle_{\text{ens}}$ is larger than $\langle E \rangle_{\text{exp}}$ indicating either too short end-to-end distances overall, or some conformations with strongly underestimated end-to-end distances. Although for there is no disagreement with CS data ($\chi^2 < 1$), the PRE-only ensemble is in worse agreement with the CS data than the TraDES RC or SAXS-only ensemble.

When the overall distribution of inter-residue distances from SAXS and the specific pattern of inter-residue distances from PRE are synthesized in one ensemble model, the transfer efficiency calculated from the ensemble, $\langle E \rangle_{\text{ens}}$, is in excellent agreement with the experimental transfer efficiency, $\langle E \rangle_{\text{exp}}$. The fit of the CS data (which were not used as a restraint for this ensemble) are also improved relative to the TraDES RC, the SAXS-only, and PRE-only ensembles. As was previously observed, generating ensembles by satisfying tertiary structure restraints seems to place some restraints on the backbone conformations[54].

Although the ensembles considered thus far are all consistent with CS data, we also calculated ensembles jointly restrained by SAXS, PRE, and CS data. However, introducing CSs (a non-distance restraint) decreases the relative weighting of distance-based restraints (SAXS and PRE) and causes a greater dispersion between $\langle E \rangle_{\text{ens}}$ for independently calculated ensembles. We therefore used a strategy of generating
ensembles jointly restrained by SAXS, PRE, and CS data and filtered them against experimental transfer efficiencies (Table 1, SAXS+PRE+CS+⟨E⟩-filter).

2.3 Integrative modelling provides a richer description of global dimensions than can be provided by SAXS or smFRET individually

To better understand the implications and advantages of combining multiple datasets we calculated global descriptions of Sic1 and pSic1 conformational ensemble dimensions (\(R_g\), \(R_{ee}\), and hydrodynamic radius \(R_h\)). SI Appendix Table S10 summarizes the global dimensions of five independently calculated ensembles with 100 conformations each (\(N_{conf} = 100\)).

The SAXS+PRE restrained ensembles have a mean \(R_{ee} = 63.6 \pm 1.1\) Å for Sic1 and \(R_{ee} = 64.7 \pm 0.7\) Å for pSic1. There is no significant discrepancy between \(R_{ee}\) inferred from smFRET using homopolymer models and the integrative approach (≤5% error). Inferences of \(R_{ee}\) using only the SAXS data overestimate the calculated mean \(R_{ee}\) by greater than 10% and depend highly on the initial conformer generation and the optimization/selection algorithm. Our approach of joint refinement/validation using SAXS, PRE and smFRET data addresses this issue. The Sic1 and pSic1 SAXS+PRE ensembles have back-calculated transfer efficiencies, \(\langle E \rangle_{ens}\), which differ by five standard deviations, while their mean \(R_{ee}\) differ by less than one standard deviation. The increased sensitivity of \(\langle E \rangle\) over \(R_{ee}\) is due to the highly non-linear \(r^6\) distance averaging, and that heterogeneous ensembles are not enforced to uniformly compact or expand to match \(\langle E \rangle_{exp}\). This demonstrates an additional advantage of integrative approaches, which use smFRET transfer efficiencies directly as restraints or validation, rather than using derived values from the data via polymer theory assumptions.

The Sic1 and pSic1 SAXS+PRE ensembles’ mean \(R_g\) (\(R_g = 29.50 \pm 0.06\) Å for Sic1 and \(R_g = 30.68 \pm 0.08\) Å for pSic1) are within 3% and 5% respectively of the model-free and the SAXS-only explicit chain estimates of \(R_g\). In contrast, the determination of \(R_g\) from smFRET strongly depends on the model used. As shown below, the calculated ensembles have smaller values of \(G\) than do homopolymers resulting in systematically underestimated \(R_g\) when the polymer-theoretic values of \(G\) are used.

The calculated hydrodynamic radius, \(R_h\), was found to be highly similar for all considered ensembles \((R_h \approx 21\) Å). The \(R_h\) of these ensembles is in excellent agreement with previously published pulsed-field gradient (PFG) diffusion NMR experiments, \(R_h = 21.5 \pm 1.1\) Å and \(R_h = 19.4 \pm 1.6\) Å for Sic1 and pSic1, respectively [34], and Fluorescence Correlation Spectroscopy (FCS) measurements \(R_h = 22 \pm 2\) Å for Sic1[55].

2.4 Analysis of the conformational behaviour of calculated ensembles beyond global dimensions

We next sought to determine descriptions of the calculated conformational ensembles which go beyond global dimensions and would facilitate comparison with polymer theory reference states, and with IDPs and unfolded states of varying sequence and chain length, \(n\). To this end, we used the fact that many aspects of homopolymer behaviour become universal, or independent of monomer identity, in the long chain (as \(n \to \infty\)) limit[56] (see below). This allowed us to clearly identify ways in which ensembles jointly restrained by SAXS and PRE data, and validated by smFRET data, deviate from homopolymer behaviour, and whether ensembles restrained only by SAXS data more resemble homopolymers, or the fully restrained ensembles.

For very long homopolymer chains, the scaling exponent \(\nu\) tends to one of only three possible limits (1/3, 1/2, 0.588), describing the poor-solvent, \(\theta\)-state, and excluded volume (EV)-limit respectively.
mers in these limits have well-defined universal values for the size ratios $G = R_{ee}^2/R_g^2$ and $\rho = R_g/R_h$, the overall shape of the ensemble, as characterized by the average asphericity $\langle A \rangle$ ($A \sim 0$ for a sphere and $A \sim 1$ for a rod), the relative variance in the end-to-end distance distribution $\Delta R_{ee} = \sqrt{\langle r_{ee}^2 \rangle - \langle r_{ee} \rangle^2}/R_{ee}$, and the relative variance in the distribution of the shape of individual conformations $\Delta A = \sqrt{\langle A^2 \rangle - \langle A \rangle^2}/\langle A \rangle$.

Table 2 summarizes the universal values expected for homopolymers in the $\theta$-state or the EV-limit, in the case of very long chains (EV and $\theta$-state $n \to \infty$) and for chains with similar length to Sic1 (EV $n = 90 – 100$).

As IDPs are finite-length heteropolymers, their apparent scaling exponents ($\nu_{app}$, see below), can take on intermediate values to these three limits. Similarly, their behaviour can deviate from the universal values expected for homopolymers. Table 2 shows the nominally universal values calculated for the experimentally-restrained ensembles. The TraDES RC, though not a homopolymer, is constructed with only excluded volume long-range interactions, and so is expected to have behaviour consistent with polymer theory predictions for an EV-limit polymers of similar chain-length (EV $n = 90 – 100$ Table 2).

![Table 2](image)

**Table 2**: Nominally universal polymer properties of the TraDES RC ensemble, SAXS-only ensemble, and SAXS+PRE ensembles

<table>
<thead>
<tr>
<th>Polymer Theory</th>
<th>$G$</th>
<th>$\rho$</th>
<th>$\langle A \rangle$</th>
<th>$\Delta A$</th>
<th>$\Delta R_{ee}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>EV ($n \to \infty$)</td>
<td>6.254</td>
<td>~1.59</td>
<td>0.431</td>
<td>0.442</td>
<td>0.374</td>
</tr>
<tr>
<td>EV ($n = 90 – 100$)</td>
<td>6.32</td>
<td>1.27–1.39</td>
<td>0.4377</td>
<td>0.437</td>
<td>-</td>
</tr>
<tr>
<td>$\theta$-state ($n \to \infty$)</td>
<td>6</td>
<td>~1.5</td>
<td>0.396</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sic1</th>
<th>TraDES RC</th>
<th>6.37</th>
<th>1.33</th>
<th>0.438</th>
<th>0.438</th>
<th>0.352</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SAXS-only</td>
<td>6.34</td>
<td>1.35</td>
<td>0.447</td>
<td>0.430</td>
<td>0.363</td>
</tr>
<tr>
<td></td>
<td>SAXS+PRE</td>
<td>4.99</td>
<td>1.33</td>
<td>0.349</td>
<td>0.461</td>
<td>0.417</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>pSic1</th>
<th>TraDES RC</th>
<th>6.35</th>
<th>1.33</th>
<th>0.438</th>
<th>0.432</th>
<th>0.366</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SAXS+PRE</td>
<td>4.83</td>
<td>1.31</td>
<td>0.361</td>
<td>0.440</td>
<td>0.388</td>
</tr>
</tbody>
</table>

* Reported values are the mean of 5 independently calculated $N_{\text{conf}} = 100$ ensembles. Table is reproduced in supplementary information with standard deviations of reported values and references for polymer theory values.

The values of $G$ for the RC and SAXS-only ensembles are indistinguishable from the expected value for a homopolymer in the EV-limit ($G \approx 6.3$). Modelling the TraDES RC using phosphorylated residues at phosphorylation sites does not change $G$, consistent with the predicted universality. In contrast, for SAXS+PRE ensembles, $G$ decreases from $G \approx 5.0$ for Sic1 to $G \approx 4.8$ for pSic1. Both values are outside the range $G_\theta = 6 \leq G \leq G_{EV} \approx 6.3$ despite the intermediate values of the apparent scaling exponents for Sic1 and pSic1 (see below).

For Sic1 and pSic1, $\rho$ is not sensitive to deviations from homopolymer statistics at long sequence separations. The value of $\rho$ remains ~1.3 for the RC, SAXS-only, and SAXS+PRE restricted ensembles, despite large changes in $R_{ee}$ and $G$. The observed $\rho$ is consistent with the range of polymer-theoretic values for a finite length EV homopolymer (EV $n = 90 – 100$ Table 2).

The Sic1 and pSic1 RC ensembles, have an average asphericity $\langle A \rangle$ very close to the polymer-theoretic value for a homopolymer in the EV-limit. The Sic1 SAXS-only ensembles are slightly more aspherical than the RC ensembles, consistent with the expected correlation between $R_g$ and $\langle A \rangle$ [21, 24]. Sic1 SAXS+PRE ensembles, however, are more spherical, with significantly lower $\langle A \rangle$, despite their larger-than-RC $R_g$. Similar to $G$, the values of $\langle A \rangle$ for the SAXS+PRE ensembles are not bound between the value predicted for the $\theta$-state and EV-limit, despite these ensembles having intermediate values of the apparent scaling exponents (see below).
The relative variance in the end-to-end distance distribution, $\Delta R_{ee}$, is close to the EV-limit value ($\Delta R_{ee}^{EV} \approx 0.37$) for the TraDES RC and Sic1 SAXS-only restrained ensembles. In contrast, $\Delta R_{ee} \approx 0.42$ for the Sic1 SAXS+PRE ensembles, which is practically identical to the $\theta$-state value. Sic1, although more compact than the RC, exhibits strong fluctuations in the end-to-end distance. Multisite phosphorylation appears to slightly reduce $\Delta R_{ee}$, although it remains above the EV-limit values.

The RC and SAXS-only ensembles have a relative variance in the distribution of shapes, $\Delta A$, similar to that of an EV-limit homopolymer, while that of the Sic1 SAXS+PRE ensemble is slightly larger. The broadness of the SAXS+PRE ensembles’ $A$ distribution stresses the fact that despite being more spherical than an EV polymer, the Sic1 ensemble contains conformations with a large distribution of shapes.

### 2.5 Internal scaling profiles and apparent scaling exponents

To extract further insights regarding the effects of combining multiple solution data types on the statistics of internal distances in the ensembles, we calculated internal scaling profiles (ISPs, Fig. 3). ISPs quantify the mean internal distances ($\langle R_{ij} \rangle = \langle r_{ij}^2 \rangle^{1/2}$) between all pairs of residues that are $|i-j|$ residues apart in the linear amino acid sequence (see Materials and Methods). The dependence of $R_{\text{app}}$ on sequence separation $|i-j|$ is often quantified by fitting to the power-law relation:

$$R_{|i-j|} = \sqrt{2l_p b} |i-j|^\nu_{\text{app}}$$  \hspace{1cm} (1)$$

where $b = 3.8 \text{ Å}$ is the distance between bonded $C_\alpha$ atoms and $l_p \approx 4 \text{ Å}$ is the persistence length. This persistence length was found to be applicable to a broad range of denatured and disordered states[5, 21, 57].

ISPs highlight important differences between ensembles. If the majority of internal distances are similar in the ISPs of two ensembles, their $R_g$ values will be similar, as $R_g = \sqrt{\frac{1}{2n} \sum_{ij} \langle r_{ij}^2 \rangle}$ [21]. However, if their spatial separations start to diverge at long sequence separations, the ensembles will have dissimilar $R_{ee}$ and $\langle E \rangle_{\text{exp}}$, when terminally labelled. This is illustrated by Fig. 3 A which shows the ISPs of the SAXS-only and SAXS+PRE Sic1 ensembles, which have similar $R_g$, but only the SAXS+PRE ensemble is consistent with the smFRET data.

Similarly, ISPs explain how $R_{ee}$ and $R_g$ can become decoupled for finite-length heteropolymers[21, 22]. Internal distances in very long homopolymers are expected to follow power-law scaling with a single global $\nu_{\text{app}}$ that defines the scaling behaviour at all sequence separations. We define the change in scaling behaviour at long sequence separations ($\nu_{\text{app}}^{\text{long}}$) relative to intermediate sequence separations ($\nu_{\text{app}}^{\text{int}}$) as $\Delta \nu_{\text{app}}^{\text{ends}} = \nu_{\text{app}}^{\text{long}} - \nu_{\text{app}}^{\text{int}}$. For long homopolymers we expect $\Delta \nu_{\text{app}}^{\text{ends}} \approx 0$. Negative (positive) values of $\Delta \nu_{\text{app}}^{\text{ends}}$ indicate ensembles with $G$ less than (greater than) predicted from $\nu_{\text{app}}$. The ISPs of SAXS-only (Fig. 3 A) and TraDES RC ensembles (Fig. 3 B&C) have $\Delta \nu_{\text{app}}^{\text{ends}} \approx 0$ consistent with the finding that these ensembles exhibit homopolymer behaviour (Table 2). In contrast, the SAXS+PRE ensembles, which are consistent with the smFRET data, have $\Delta \nu_{\text{app}}^{\text{ends}} < 0$, consistent with lower than expected $G$ and $\langle A \rangle$[21].

To rigorously quantify deviations homopolymer statistics, we fit five independently calculated ensembles with 100 conformations ($N_{\text{conf}} = 100$, Table 3). In an intermediate regime (15 $\leq |i-j| \leq 51$), Sic1 and pSic1 SAXS+PRE ensembles have a scaling exponent $\nu_{\text{app}}^{\text{int}} \approx 0.53$, which suggests that at these scales, the physiological buffer is a marginally good solvent. At longer sequence separations (51 $< |i-j| \leq n_{\text{res}} - 5$), the ensembles show behaviour which is closer to the poor solvent scaling regime $\nu_{\text{long}} \approx 0.3$. We performed a paired t-test on the five $N_{\text{conf}} = 100$ ensembles to determine if the differences $\Delta \nu_{\text{app}}^{\text{ends}}$ come from a distribution with zero mean (Table 3). The deviations from homopolymer statistics (i.e.,
\( \Delta \nu_{\text{app}}^{\text{ends}} \neq 0 \) are statistically significant (p-value \(< 0.01\)) for the SAXS+PRE ensembles but not for the TraDES RC ensembles (p-value \(\approx 0.14\)). The SAXS-only ensemble has weaker evidence for deviations from ideal homopolymer statistics (p-value \(\approx 0.03\)). Ensembles restrained only by SAXS data are more similar to EV homopolymers, than to the fully experimentally restrained ensembles.

The Sic1 and pSic1 SAXS+PRE ISPs in Fig. 3 A are consistent with NMR, SAXS and smFRET data. We therefore consider these ISPs as the benchmarks in determining the scaling behaviour of the Sic1 ensemble and compare it to recently published methods which infer scaling behaviour from only SAXS[22, 29] (SI Appendix, Table S6), or only smFRET data (\(\nu_{\text{app}} \approx 0.52\) for Sic1 and \(\nu_{\text{app}} \approx 0.54\) for pSic1)[30]. Due to the terminal labelling positions and \(\Delta \nu_{\text{app}}^{\text{ends}} < 0\), the \(\nu_{\text{app}}\) inferred from smFRET is less than that inferred from SAXS, though neither give as complete a picture of the scaling behaviour as the ISPs from integrative modelling. Using a heterpolymer MFF analysis, which allows for deviations in power-law scaling at long sequence separations (MFF-het3, SI Appendix, Table S6), gave similar results to separately fitting ISPs at intermediate and long-sequence separations (\(\nu_{\text{app}} = 0.56 \pm 0.01\) and \(\Delta \nu_{\text{app}}^{\text{ends}} = -0.17 \pm 0.08\) for Sic1). However, in the absence of additional measurements, it is not clear when to prefer this model over the equally well-fit but more parsimonious homopolymer MFF models.

**Figure 3:** (A) \(N_{\text{conf}} = 500\) Sic1 SAXS+PRE ensembles (circles) and Sic1 SAXS-only ensembles (squares) with fits to intermediate (dashed) and long (solid) sequence separations. (B) \(N_{\text{conf}} = 500\) Sic1 SAXS+PRE ensembles (circles) and Sic1 TraDES RC (squares) with fits to intermediate (dashed) and long (solid) sequence separations. (C) \(N_{\text{conf}} = 500\) pSic1 SAXS+PRE ensembles (black) and pSic1 TraDES RC (red) with fits to intermediate (dashed) and long (solid) sequence separations. For visualization, every fifth data point is shown.

**Table 3:** Fitting results for the TraDES RC ensemble, SAXS-only ensemble, and SAXS+PRE ensembles ISPs

<table>
<thead>
<tr>
<th></th>
<th>TraDES RCb</th>
<th>Sic1 SAXS-only</th>
<th>Sic1 SAXS+PRE</th>
<th>pSic1 SAXS+PRE</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\nu_{\text{app}}) (fixed (l_p = 4) Å)</td>
<td>0.570</td>
<td>0.585</td>
<td>0.579</td>
<td>0.588</td>
</tr>
<tr>
<td>(\nu_{\text{app}}^{\text{int}})</td>
<td>0.566</td>
<td>0.583</td>
<td>0.527</td>
<td>0.524</td>
</tr>
<tr>
<td>(\nu_{\text{app}}^{\text{long}})</td>
<td>0.51</td>
<td>0.51</td>
<td>0.30</td>
<td>0.31</td>
</tr>
<tr>
<td>(\Delta \nu_{\text{app}}^{\text{ends}})</td>
<td>-0.06</td>
<td>-0.07</td>
<td>-0.23</td>
<td>-0.21</td>
</tr>
<tr>
<td>Paired t-test p-valuec</td>
<td>0.143</td>
<td>0.027</td>
<td>1.6 \times 10^{-3}</td>
<td>5.1 \times 10^{-4}</td>
</tr>
</tbody>
</table>

* Table results are the mean results from fitting 5 \(N_{\text{conf}} = 100\) ensembles. Standard deviation of the mean for \(\nu_{\text{app}}^{\text{long}}\) and \(\nu_{\text{app}}^{\text{int}}\) is \(\approx 0.005\) and for \(\nu_{\text{app}}^{\text{ends}}\) is \(\approx 0.03\). See Materials and Methods for additional details.

* Sic1 TraDES RC and pSic1 TraDES RC result in nearly identical fits.

* Paired t-test for \(\nu_{\text{app}}^{\text{int}}\) and \(\nu_{\text{app}}^{\text{long}}\) differences.
2.6 Two dimensional scaling maps reveal regional biases for expansion and compaction

To better describe the heteropolymeric nature of Sic1, a normalized two-dimensional (2D) scaling map was constructed (Fig. 4). In the first step, the ensemble-averaged distances between the C\textsubscript{\(\alpha\)} atoms of every unique pair of residues in the sequence is calculated for the experimentally-restrained ensemble (\(\langle r^2_{ij} \rangle_{\text{ens}}\)), and for the respective TraDES RC ensemble (\(\langle r^2_{ij} \rangle_{\text{RC}}\)). Experimentally-restrained distances are normalized by the RC distances and displayed as a 2D scaling map.

The normalized 2D scaling map for Sic1 (Fig. 4 A) displays regional biases for expansion (\(\alpha_{ij} > 1\)) and compaction (\(\alpha_{ij} < 1\)). Short internal distances \(|i-j| \lesssim 45\) show expansion relative to the RC, while \(|i-j| \gtrsim 60\) show compaction. The expansion, however, is heterogeneous. For example, the \(\sim 40\) residue N-terminal region is significantly more expanded than the \(\sim 40\) residue C-terminal region. Similar distinctions between the RC and pSic1 ensembles were observed (Fig. 4 B).

![Figure 4](image)

Figure 4: (A) Sic1 2D scaling map \(\alpha_{ij} = \langle r^2_{ij} \rangle_{\text{ens}}^{1/2}/\langle r^2_{ij} \rangle_{\text{RC}}^{1/2}\) using the Sic1 (SAXS+PRE) \(N_{\text{conf}} = 500\) and the Sic1 \(N_{\text{conf}} = 500\) TraDES RC ensemble. (B) pSic1 2D scaling map \(\alpha_{ij} = \langle r^2_{ij} \rangle_{\text{ens}}^{1/2}/\langle r^2_{ij} \rangle_{\text{RC}}^{1/2}\) using the pSic1 (SAXS+PRE) \(N_{\text{conf}} = 500\) and the pSic1 \(N_{\text{conf}} = 500\) TraDES RC ensemble. (C) pSic1 normalized by Sic1 dimensions.

To compare Sic1 and pSic1 ensembles, the pSic1 ensemble was normalized by the Sic1 ensemble, (Fig. 4 C). This map describes the heterogeneous modulation of Sic1 upon multisite phosphorylation. Sic1 expansion upon phosphorylation has been attributed to transient tertiary contacts involving non-phosphorylated CPDs that are lost or weakened upon phosphorylation[33]. In our ensembles, expansion is also seen to be clustered around CPD sites, particularly those of the C-terminus. Expansion is also seen in the vicinity of Y14, previously implicated in tertiary interactions with CPDs[34] (see below).

2.7 Y14A mutation and phosphorylation disrupt tertiary contacts in Sic1

An intriguing possibility is that specific tertiary contacts, involving pi-pi[58] and cation-pi interactions[59], lead to compaction in Sic1. The Sic1 sequence is 23% residues with side chains containing pi-groups (F,N,Q,R, and Y), and 52% residues with relatively exposed pi-groups in peptide backbone amide groups (G,P,S, and T) [58]. In particular, PRE effects link CPDs with Y14 and \(^{15}\)N relaxation experiments on Sic1 identified maxima in the \(R^2\) rates near Y14[34]. Furthermore, the substitution Y14A led to an expansion in \(R_h\) of \(\sim 20\%\) in pSic1 [34]. We hypothesized that if Y14 engages in specific pi-pi and cation-pi interactions\(^1\) throughout the chain, then removing its pi-character by mutation to alanine will disrupt these interactions, leading to larger \(R_{ee}\) and lower \(\langle E \rangle_{\text{exp}}\). On the Kyte-Doolittle scale[60] this mutation increases the hydrophobicity

\(^1\)The Sic1 sequence has 6 K and 5 R residues. The predominantly 6- and 7-fold phosphorylated Sic1 studied here therefore has a net charge (excluding the dyes) of -1 to -3 assuming each phosphate group contribute -2 charge.
\(H_Y = 0.36 \rightarrow H_A = 0.7\) suggesting that expansion would not result from reduced hydrophobicity.

Figure 5: Y14A mutation and phosphorylation results in a shift to lower \(\langle E \rangle_{\text{exp}}\) (more expanded conformations). Each histogram is normalized so that each Gaussian fit has a maximum of one.

We performed smFRET experiments for the Y14A mutants of Sic1 and pSic1 (Fig. 5 and SI Appendix, Table S9). Y14A mutation decreases Sic1 \(\langle E \rangle_{\text{exp}}\) by approximately 7\% (ca. 0.42 to 0.40, a small but reproducible shift). Similarly, Y14A mutation decreases pSic1 \(\langle E \rangle_{\text{exp}}\) by approximately 8\% (ca. 0.36 to 0.33). These results are consistent with chain expansion driven by the disruption of interaction with the pi-group of the tyrosine Y14. The cumulative effect of Y14A mutation and phosphorylation on Sic1 is to decrease \(\langle E \rangle_{\text{exp}}\) by approximately 22\%, such that \(\langle E \rangle_{\text{exp}} = 0.33 \pm 0.02\) is similar to that of the Sic1 TraDES RC \((\langle E \rangle_{\text{ens}} = 0.30 \pm 0.01)\).

3 Discussion

We generated SAXS and smFRET data on Sic1 and pSic1, and resolved their apparently discrepant inferences by joint refinement by the SAXS and PRE data. The ensembles restrained by SAXS and PRE data are, in addition, consistent with the smFRET data, chemical shift data, and hydrodynamic data (PFG-NMR and FCS). We used smFRET transfer efficiencies directly as validation, rather than using derived values from the data via polymer theory assumptions. Our final ensembles of Sic1 can be examined with a high degree of confidence given their agreement with a diverse set experimental data acting as both restraints and validation.

We therefore explored the differences between ensembles restrained by different data types, and how they compare to homopolymer reference states, by calculating global descriptions of Sic1 and pSic1 conformational ensemble dimensions \((R_g, R_{ee}, R_h)\), their size ratios \((G\text{ and }\rho)\), overall shape \((\langle A \rangle)\) and the relative variances of their end-to-end distance and shape distributions \((\Delta R_{ee}\text{ and }\Delta A)\). To extract further insights into their length-scale and regional biases for expansion and compaction, we calculated internal scaling profiles and 2D scaling maps. We examined relationships between average global polymeric descriptions and higher-moments of their distributions, which are prerequisite for a complete description of sequence-to-ensemble and ensemble-to-function relationships of IDPs [32].

The picture that emerges when the entirety of the experimental data on Sic1 and pSic1 is considered, is that their conformational ensembles cannot be described by statistics derived for infinitely long homopolymers. Although this is unsurprising, given that Sic1 and pSic1 are finite-length heteropolymers, ensembles restrained only by the SAXS data are congruent with the set of homopolymer descriptions and scaling
relationships for EV homopolymers. Neither the SAXS nor smFRET data, individually, suggest significant deviations from homopolymer statistics. Our results therefore provide a strong impetus for integrative modelling approaches over homopolymer approaches whenever multiple data types exist.

We emphasize that the SAXS+PRE ensembles were not constructed by reweighting or selecting ensembles specifically to achieve agreement with \( \langle E \rangle_{\exp} \). In our approach, it was not guaranteed \emph{a priori} that \( \langle E \rangle_{\text{ens}} \) would match \( \langle E \rangle_{\exp} \), especially if either the introduction of PRE spin labels or smFRET fluorophores had perturbed the IDP ensemble. If negative values of \( \Delta \nu_{\text{ends}}^{\text{app}} \) are common in IDPs and unfolded proteins under refolding conditions, smFRET on terminally labelled samples will infer smaller \( \nu_{\text{app}} \) relative to SAXS. Since \( \Delta \nu_{\text{ends}}^{\text{app}} \) appears to be undetectable without integrative modelling, the effect would be qualitatively similar to fluorophore-protein interactions, and like fluorophore-protein interactions it would disappear in high concentrations of denaturant where \( \Delta \nu_{\text{ends}}^{\text{app}} \approx 0 \)[21, 22]. Deciding between fluorophore-interactions and heteropolymer effects requires an integrative modelling approach.

To fully understand the practical utility of different restraint types for characterizing types of structure in IDPs will require a more rigorous approach for scoring the probability of an ensemble on the basis of its agreement with diverse experimental data using Bayesian procedures[61–63]. For example, Lincoff and co-workers recently developed a Bayesian scoring formalism, the extended Experimental Inferential Structure Determination (X-EISD) method, to calculate the most probable ensembles for the drk SH3 unfolded state domain [64]. Using this method, they also found that FRET and PRE provide strong discriminatory power in determining the most probable ensemble.

### 3.1 Conformation-to-function relationships

For soluble post-translationally modified IDPs, approximately good-solvent scaling may be unsurprising. The balance between chain-chain and chain-solvent interactions is a driving force for aggregation[65] and phase separation[66, 67], and polymer theory predicts that proteins with overall good-solvent scaling in native-like conditions should remain soluble, while chains with poor-solvent scaling are expected to undergo aggregation or phase separation. At short-to-intermediate sequence separations, good-solvent scaling provides read/write access of substrate motifs to modifying enzymes (e.g., phosphorylation and ubiquitination for Sic1).

Good-solvent scaling also confers advantages specifically to fuzzy or dynamic complexes as internal friction increases with increasing chain compaction[68]. Low internal friction and fast chain reconfigurations are therefore expected for short-to-intermediate separations. In a kinetic model of ultrasensitive binding, fast reconfiguration dynamics provides more opportunities for unbound CPDs to (re)bind before pSic1 diffuses out of proximity of Cdc4[47, 48, 69]. In the polyelectrostatic model, fast reconfiguration dynamics facilitates pSic1’s dynamic interactions with Cdc4 through electrostatic averaging effects[34, 46].

The crossover to poor-solvent scaling at long sequence separations, \( G < G_\theta < G_{EV} \), and \( \langle A \rangle < \langle A \rangle_{\theta} < \langle A \rangle_{EV} \), imply that unbound CPDs that are sequence-distant from a bound CPD are on average closer to the WD40 binding pocket than they would be for an EV-chain, thus decreasing the solvent screening of electrostatic interactions. A prediction of the polyelectrostatic model is that decreasing the distance between the binding pocket and the overall mean-field charge distribution of Sic1, leads to sharper transitions in the fraction bound with respect to the number of phosphorylations[46]. In the kinetic model, these deviations from EV-statistics increase the effective concentration of CPDs in the vicinity of the binding pocket, which may increase the probability for any CPD to rebind before diffusive exit.

Although Sic1 ensembles are more spherical than EV-limit ensembles \( \langle A \rangle < \langle A \rangle_{\theta} < \langle A \rangle_{EV} \), the relative fluctuations in shape exceed those of the EV-limit ensembles (Table 2). Similarly, the relative fluctuations
in $R_{ee} (\Delta R_{ee})$ for these ensembles exceeds the expected fluctuations not only for an EV-limit polymer, but also those of the $\theta$-state (Table 2). Large amplitude fluctuations in the shape and size of Sic1, effectively and rapidly sampling many different conformations, could allow CPDs in Sic1 to rapidly sample either the primary or secondary WD40 binding pocket, before the two proteins diffuse away. These fluctuations could also facilitate electrostatic averaging, permitting a mean-field treatment as assumed in the polyelectrostatic model.

4 Conclusions

Our work provides a high-resolution description of the conformational ensembles of Sic1 and pSic1 in physiological conditions. Our calculated ensembles are consistent with experimental data reporting on a wide range of spatial and sequence separation scales: local/secondary structure (CSs), non-local/tertiary structure (PRE and smFRET) and global/molecular size and shape (SAXS, PFG-NMR and FCS). Our results show that there are clear advantages of combining multiple datasets and that quantitative polymer-physics-based characterization of experimentally-restrained ensembles could be used to implement a rigorous taxonomy for the description and classification of IDPs as heteropolymers. The chain length independence of many of these properties facilitates comparison between different IDPs and unfolded states of proteins.

Our results suggest that for Sic1 and our dye pair, discrepant inferences between SAXS and smFRET cannot a priori be assumed to arise from “fluorophore-interactions.” The impact of the fluorophores (or spin-labels) will of course depend on the physicochemical properties of the specific IDP sequence and the fluorophores (or spin-labels) used. Robustness to perturbation (e.g., labels or phosphorylation) may be built into Sic1’s sequence via its patterning of charged and proline residues, as observed for other substrates of proline directed kinases (e.g., Ash1 [8]). Further ensemble modelling of IDPs will reveal to what extent robustness to labelling, and deviations from homopolymer statistics, are general to IDPs. In this regard, an integrative use of multiple experiments probing disparate scales, computational modelling, and polymer physics, will provide valuable insights into IDPs/unfolded states and their biological functions.

5 Materials and Methods

5.1 Sic1 samples

The Sic1 N-terminal IDP region (1-90, henceforth Sic1) was expressed recombinantly as a Glutathione S-transferase (GST) fusion protein in Escherichia coli BL21 (DE3) codon plus cells and purified using glutathione-Sepharose affinity chromatography and cation-exchange chromatography. The correct molecular mass of the purified protein was verified by electrospray ionization mass spectrometry (ESI-MS).

A double cysteine variant of Sic1 (-1C-T90C) for smFRET experiments was generated via site directed mutagenesis from a single-cysteine mutant produced previously for PRE measurements[33, 34]. This construct was purified as above and the correct molecular mass of the purified protein was verified by ESI-MS. A Y14A mutant Sic1 (-1C-T90C-Y14A) was generated via site directed mutagenesis from the aforementioned double-cysteine mutant and was expressed, purified, and characterized using the same protocol.

The Sic1 smFRET construct was labelled stochastically with Alexa Fluor 488 C5 Maleimide (ThermoFisher Scientific, Invitrogen, A10254) and Alexa Fluor 647 C2 Maleimide (ThermoFisher Scientific, Invitrogen, A20347). After labelling with Alexa Fluor 647, cation-exchange chromatography was used to separate species with a single acceptor label, from doubly acceptor labelled and unlabelled species. The single-labelled species sample was then labelled with Alexa Fluor 488 and cation-exchange chromatography was used to separate doubly heterolabelled from acceptor only species. The correct mass of the doubly labelled sample was confirmed by mass spectrometry. The final FRET labelled sample was concentrated and buffer exchanged into PBS buffer pH 7.4 with 3 M GdmCl, 2 mM DTT and stored at -80 °C.
Phosphorylated samples were prepared by treatment of Sic1 with Cyclin A/Cdk2 (prepared according to Huang et al., [70]) at a kinase:Sic1 ratio of 1:100 in the presence of 50 fold excess of ATP and 2.5 mM MgCl₂ overnight at 30 °C. The yield of phosphorylation reaction was determined by ESI-MS. Under these conditions the dominant species are 6- and 7-fold phosphorylated Sic1 (10195 Da and 10274 Da respectively) with a small fraction of 5-fold phosphorylated Sic1. After phosphorylation, the samples were buffer exchanged into PBS buffer pH 7.4 with 3 M GdmCl to prevent aggregation, denature kinase, and denature any phosphatases which may have inadvertently entered the solution. The samples were kept on ice in 4°C and measured within 24 hours.

Additional details regarding protein expression, purification and labelling are available in the supplementary information.

5.2 Single-molecule fluorescence

Single-molecule fluorescence experiments were performed on a custom-built multiparameter confocal microscope with microsecond alternating laser excitation. This instrumentation allows the simultaneous detection of the intensity, anisotropy, lifetime, and spectral properties of individual molecules and for the selection of fluorescence bursts in which both dyes are present and photophysically active.

Immediately prior to measurement samples were diluted to ~50 pM in either (i) PBS buffer: 10 mM sodium phosphate and 140 mM NaCl pH 7.0, 1 mM EDTA (to replicate NMR measurement buffer of Ref [33]) or (ii) Tris buffer: 50 mM Tris and 150 mM NaCl, pH 7.5. (to replicate SAXS measurement buffer). No difference in $\langle E \rangle_{exp}$ was detected when comparing buffer conditions and results are shown for Tris buffer conditions.

The acquired data were subjected to multiparameter fluorescence analysis[71, 72] and ALEX filtering[73]. The burst search was performed using an All Photon Burst Search (APBS)[74, 75] with $M = 10$, $T = 500 \mu$s and $L = 50$. Transfer efficiencies were determined burst-wise and corrected for differences in the quantum yields of the dyes and detection efficiencies, as described in further detail in the SI Appendix.

The Förster radius $R_0$ was calculated assuming a relative dipole orientation factor $\kappa^2 = 2/3$ and the refractive index of water $n = 1.33$. The assumption of $\kappa^2 = 2/3$ is supported by subpopulation-specific steady-state anisotropies for the donor in the presence of the acceptor $r_{DA}$ (SI Appendix, Table S1). The overlap integral $J$ was measured for each sample and found not to change upon phosphorylation or Y14A mutation. The minimal variation in donor-only lifetimes $r_{DA}$ suggested minimal variation in the donor-quantum yield $\phi_D$. $R_0$ was therefore calculated to be $R_0 = 52.2 \pm 1.1 \text{ Å}$ for all samples, and variation between samples within this uncertainty.

We estimate the precision for $\langle E \rangle_{exp}$ to be ca. 0.005 (for measurements performed on the same day, with approximately equal sample dependent calibration factors). We estimate the accuracy of $\langle E \rangle_{exp}, \sigma_{E,exp}$, to be ca. 0.02 (due to uncertainty in the instrumental and sample dependent calibration factors). Further details about the instrumentation, photoprotection, laser excitations, burst detection, filtering and multiparameter fluorescence analysis can be found in the SI Appendix.

5.3 Small-angle X-ray scattering

Small angle X-ray scattering data were collected at beamline 12-ID-B at the Argonne National Laboratory Advanced Photon Source. Protein samples were freshly prepared using size exclusion chromatography (GE Life Sciences, Superdex 75 10/300 GL) in a buffer containing 50 mM Tris pH 7.5, 150 mM NaCl, 5 mM DTT, and 2 mM TCEP. Fractions were loaded immediately after elution without further manipulation. Buffer collected one column volume after protein elution from the column was used to record buffer data before and after each protein sample. SAXS data were acquired manually; protein samples were loaded, then gently refreshed with a syringe pump to prevent x-ray damage. A Pilatus 2M detector provided q-range coverage from 0.015 Å⁻¹ to 1.0 Å⁻¹. Wide-angle x-ray scattering data were acquired with a Pilatus 300k detector and had a q range of 0.93 – 2.9 Å⁻¹. Calibration of the q-range calibration was performed with a silver behenate sample. Twenty sequential images were collected with 1 sec exposure time per image with each detector. Data were inspected for anomalous exposures and mean buffer data were subtracted from sample data using the WAXS water peak at $q \sim 1.9 \text{ Å}^{-1}$ as a subtraction control. Details about the SAXS data analysis can be found in the SI Appendix.

5.4 ENSEMBLE

ENSEMBLE 2.1 [35] was used to determine a subset of conformations from an initial pool of conformers created by the statistical coil generator TraDES[49, 50]. For restraining with SAXS and PRE data, both modules were given rank 1. When SAXS, PRE and CS modules were used, PRE and SAXS were given rank 1 and CSs were given rank 2. All other ENSEMBLE parameters were left at their default values.
To achieve a balance between the concerns of over-fitting (under-restraining) and under-fitting (over-restraining) we performed multiple independent ENSEMBLE calculations with 100 conformers, \( N_{\text{conf}} = 100 \), as suggested by Ref [54], and averaged the results from independent ensemble calculation or combined them to form ensembles with larger numbers of conformers (e.g., \( N_{\text{conf}} = 500 \)). Structural features resulting from over-fitting (fitting the “noise” in the experimental data) should be averaged out in independent ENSEMBLE calculations, while rare states which are conserved in independently calculated ensembles (and thus have evidence from the data) should accumulate in weight when the ensembles are combined[54]. The agreement with experimental data and polymer-theory based ensemble descriptions were highly similar for repeated independently calculated ensembles (see SI Appendix Table S10-11). This supports their averaging or consolidation into one larger ensemble. Similarly, Lincoff and coworkers demonstrated overall convergence for \( N_{\text{conf}} = 100 \) ensembles of drk SH3 unfolded state domain[64].

NMR data was obtained from BMRB accession numbers 16657 (Sic1) and 16659 (pSic1)[33]. A total of 413 PRE restraints were used with a typical conservative upper- and lower-bound on PRE distance restraints of ±5 Å[53, 76]. This tolerance was used in computing the \( \chi^2 \) metric for the PRE data. CSs were back-calculated using the SHIFTX calculator[77] and a total of 61 \( C_\alpha \) CSs and 56 \( C_\beta \) CSs were used. The CS \( \chi^2 \) metric was computed using the experimental uncertainty \( \sigma_{\text{res}} = \pm 0.4 \) ppm and the uncertainty in the SHIFTX calculator \( (\sigma_{\text{SHIFTX}} = 0.98 \) ppm for \( C_\alpha \) CSs and \( \sigma_{\text{SHIFTX}} = 1.10 \) ppm for \( C_\beta \) CSs[77]).

CRYSOL[37] with default solvation parameters was used to predict the solution scattering from individual structures from their atomic coordinates. A total of 235 data points from \( q = 0.02 \) to \( q = 0.254 \) Å\(^{-1}\) were used in SAXS restrained ensembles. The SAXS \( \chi^2 \) metric was computed using the experimental uncertainty in each data point. In principle, the calculation of \( \chi^2 \) should also include an uncertainty in the SAXS back-calculation (in particular due to uncertainty in modelling the solvation shells of IDPs[78]).

Accessible volume (AV) simulations[38, 39] were used to predict the sterically accessible space of the dye attached to each conformation via its flexible linker. These calculations were performed using the “FRET-restrained positions and screening” (FPS) software[38] provided by the Seidel group. The mean distance between dyes for conformer \( k \), \( \langle r_{DA} \rangle_k \), is used to calculate the per-conformer FRET efficiency \( e_k (\langle r_{DA} \rangle_k ; R_0) \) (Figure 1D). As chain reconfigurations are much faster than the averaging time in smFRET, the smFRET experiment measures \( P(E) \), the probability of observing a burst with efficiency \( E \), rather than the probability distribution of per-conformer FRET efficiencies \( p(e) \) (Figure 1E-F). Ensembles are therefore evaluated by the discrepancy \( \langle E \rangle_{\text{exp}} - \langle E \rangle_{\text{ens}} \) (Figure 1F). The uncertainty in \( (\langle E \rangle_{\text{ens}}, \sigma_{E,\text{ens}}) \), is ca. 0.01, primarily due to uncertainty in \( R_0 \); a similar value was obtained by Lincoff and coworkers[64]. Differences \( |\langle E \rangle_{\text{exp}} - \langle E \rangle_{\text{ens}}| \leq \sqrt{\sigma_{E,\text{exp}}^2 + \sigma_{E,\text{ens}}^2} \approx 0.02 \) indicate no disagreement between back-calculated and experimental mean transfer efficiencies. A comprehensive description of the ENSEMBLE calculations, restraints and back-calculations can be found in the SI Appendix.

5.5 Polymer scaling analysis

The distance \( R_{i\rightarrow j}^{\text{app}} = \left\langle r_{ij}^2 \right\rangle_{\text{ens}} \) between \( C_\alpha \) atoms is an average first over all pairs of residues that are separated by \( |i - j| \) residues, and then over all conformations in the ensemble. The apparent scaling exponent \( \nu_{\text{app}} \) was estimated by fitting an ISP calculated for each \( N_{\text{conf}} = 100 \) ensemble to the following expression:

\[
\ln \left( R_{i\rightarrow j}^{\text{app}} \right) = \nu_{\text{app}} \ln (|i - j|) + A_0
\]

Eq. 2 is derived for homopolymers in the infinitely long chain limit. Following Peran and coworkers[31], for finite-length chains, a lower bound of \( |i - j| > 15 \) was used to exclude deviations from infinitely long chain scaling behaviour at short sequence-separations and an upper bound of \( |i - j| < n_{\text{res}} - 5 \) was used to exclude deviations due to “dangling ends.” With these restrictions, finite-length homopolymers are expected to be well fit by Eq. 2. Evenly spaced points in log-log space were used during fitting. Fitting the entire \( 15 < |i - j| < n_{\text{res}} - 5 \) range was used to obtain \( \nu_{\text{app}} \). \( A_0 \) was either fixed at \( \log(5.51) \) (\( \ell_p = 4 \) Å) or left as a free fitting parameter.

To test for differences in scaling behaviour at intermediate and long sequence-separations, the \( 15 < |i - j| < n_{\text{res}} - 5 \) range was evenly divided into intermediate \( \nu_{\text{int}}^{\text{app}} \) (15 \( \leq |i - j| \leq 51 \)) and long \( \nu_{\text{app}}^{\text{long}} \) regimes (51 \( < |i - j| < n_{\text{res}} - 5 \)). Homopolymers are expected to have \( \nu_{\text{app}}^{\text{long}} \approx \nu_{\text{app}}^{\text{int}} \). A paired t-test was performed in MATLAB R2018b using five \( N_{\text{conf}} = 100 \) \( \nu_{\text{app}}^{\text{long}} \) and \( \nu_{\text{app}}^{\text{int}} \) estimates, to test whether the set of \( \Delta \nu_{\text{ends}} = \nu_{\text{app}}^{\text{long}} - \nu_{\text{app}}^{\text{int}} \), come from a normal distribution with mean equal to zero and unknown variance. Tests for normality (Jarque–Bera, Anderson-Darling, Lilliefors) were performed in MATLAB 2018b (\( p > 0.4 \)). Table 3 reports the mean difference for the five ensembles.
6 Associated content

6.1 Supporting information

Extended description of smFRET and SAXS experiment/analysis, ENSEMBLE methods, and additional tables

7 Author information

7.1 Corresponding Authors

* Email: gregory.gomes@mail.utoronto.ca

* Email: claudiu.gradinaru@utoronto.ca

7.2 Author contributions

G.G. and C.G. conceived the project; G.G. designed, performed and analysed smFRET experiments; E.M., G.G., and T.M. performed and analysed SAXS experiments; M.K., J.F.K., and T.H.G. guided the implementation of the ENSEMBLE analysis; G.G. performed simulations and calculated ensembles; G.G. and C.G. wrote the manuscript; G.G., T.M., T.H.G., J.F.K., and C.G. contributed insights, discussed results and interpretations, and edited the manuscript.

7.3 Notes

The authors declare no competing financial interests

8 Acknowledgements

G.N.G., M.K., J.F.-K., and T.H.G thank the National Institutes of Health for support under Grant 5R01GM127627-03. J.F.-K. thanks the Natural Sciences and Engineering Research Council of Canada for support under RGPIN-2016-06718 Fund 490974. C.C. thanks the Natural Sciences and Engineering Research Council of Canada for support under RGPIN 2017–06030. T.M. was supported by funding from St. Jude Children’s Research and the American Lebanese Syrian Associated Charities. We thank S. Chakravarthy, J. Hopkins and all BioCAT beamline staff at the Advanced Photon Source for assistance with SAXS measurements. Use of the Advanced Photon Source was supported by the U.S. Department of Energy under contract DE-AC02-06CH11357.

References


Conformational ensembles

Global polymeric descriptions

Hypotheses about biological function

NMR+SAXS restraints

smFRET validation

Figure 6: For Table of Contents only