


 Cite this: *RSC Adv.*, 2023, **13**, 10182

The influence of random-coil chemical shifts on the assessment of structural propensities in folded proteins and IDPs†

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In studying secondary structural propensities of proteins by nuclear magnetic resonance (NMR) spectroscopy, secondary chemical shifts (SCSs) serve as the primary atomic scale observables. For SCS calculation, the selection of an appropriate random coil chemical shift (RCCS) dataset is a crucial step, especially when investigating intrinsically disordered proteins (IDPs). The scientific literature is abundant in such datasets, however, the effect of choosing one over all the others in a concrete application has not yet been studied thoroughly and systematically. Hereby, we review the available RCCS prediction methods and to compare them, we conduct statistical inference by means of the nonparametric sum of ranking differences and comparison of ranks to random numbers (SRD-CRRN) method. We try to find the RCCS predictors best representing the general consensus regarding secondary structural propensities. The existence and the magnitude of resulting differences on secondary structure determination under varying sample conditions (temperature, pH) are demonstrated and discussed for globular proteins and especially IDPs.

Received 13th February 2023

Accepted 15th March 2023

DOI: 10.1039/d3ra00977g

rsc.li/rsc-advances

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† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d3ra00977g>

1. Introduction

NMR spectroscopy is one of the few methods that can provide secondary structural information for proteins at atomic level resolution. In this respect, a crucial parameter is the chemical shift (CS) which reports on the local chemical environment of



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methods for global and local characterization of biomolecules. One main topic is the investigation of intrinsically disordered proteins/protein regions, regarding the assessment of secondary structural propensities, proline conformations, protein–protein interactions. Further topics include translational diffusion studies of folded proteins and IDPs, investigation of bicelle systems and novel drug delivery peptides.



the corresponding atom. The first step of any NMR investigation is spectral assignment and as a result, chemical shift information is obtained for various atom types. Extraction of structural information from chemical shifts is a well-established practice.^{1–10} The method relies on calculating the secondary chemical shift (SCS), defined as the difference between the measured chemical shift and the appropriate random coil chemical shift (RCCS) value:

$$\text{SCS}_i^A = \delta_{i,\text{measured}}^A - \text{RCCS}_i^A \quad (1)$$

where i indicates the position in the amino acid sequence and A indicates the atom-type which the SCS corresponds to. RCCS values are generally available for H^N , H^α , N , C^α and C' atoms of the peptide backbone and the C^β atom of the side chain. The corresponding SCSs can be used to differentiate regions with helical, extended and random-coil like secondary structure, that proved to be useful in the characterization of folded proteins, while in the study of intrinsically disordered proteins and protein regions (IDPs/IDRs) defining the secondary structural propensities is of utmost importance.

Many IDPs/IDRs have been shown to play important biological roles^{11–14} and this created the need to study and assess their physical, chemical and biological behavior. An important goal is to make IDPs pharmaceutical targets.^{15–24} In this respect – despite its inefficiency – still the earlier sequence–structure–function paradigm is utilized as a starting point of the characterization.²⁵ This means, that investigation is focused on finding remnants of structural features. Even though IDPs are generally disordered, they can exhibit inherent structural preferences²⁶ that are referred to as secondary structural propensity, residual structure, transient structure. Apart from an inclination toward the well-known, another reason for identifying regions with structural propensities is that such regions are usually the ones involved in protein/protein or protein/membrane interactions. Typical sequential regions with detectable structural propensities are the so-called preformed interaction prone fragments, preformed structural motifs and short linear motifs which are generally crucial for the function of IDPs/IDRs.^{27–31} Thus, the regions of modest structural propensity are also expected to be key to understanding interactions and achieving the druggability of IDPs. For this, an efficient experimental characterization of these regions is necessary, which most tools of protein research are unable to accomplish. Due to the high flexibility of IDPs,³² attempts *via* classical methods, based on a rather rigid three-dimensional chemical structure, are inadequate.³³ The necessity of describing IDPs in terms of structural ensembles instead of single structures especially calls for multiple sources of experimental data³⁴ further increasing the importance of NMR. Consequently, the correct interpretation of the SCS is necessary.

A typical evaluation of the SCS values calculated based on eqn (1) is *via* the graphical representation as a function of the amino acid sequence. The positive or negative sign indicates the type of secondary structural propensity, while the amplitude shows the strength of this propensity. Data can also be interpreted indirectly *via* calculating some function of the SCSs, that

can be (i) the probability of the presence of different secondary structural elements,³⁵ (ii) the so-called CheZOD Z-score,³⁶ (iii) the secondary structure propensity score (SSP),³⁷ (iv) the neighbor-corrected structural propensity score,³⁸ (v) the chemical shift index (CSI),^{39–42} (vi) the random coil index (RCI)^{43–45} and (vii) probability-based secondary structure identification (PSSI).⁴⁶ Lately, attempts have been made to base disorder prediction exclusively on Z-score values, and thus exclusively on SCSs.^{47,48} Moreover, SCSs are used in the software such as SHIFTX,⁴⁹ NMRView⁵⁰ PESCADOR⁵¹ and DIPEND.⁵² The existence of such advanced methods indicates the importance of calculating SCS values appropriately. However, they are ambiguous quantities, where the ambiguity arises already from the definition (see eqn (1)), by the involvement of RCCSs. The real value of the RCCS of a given atom, for a given amino acid, under given experimental conditions is not well-defined. This is proved by the number of RCCS calculation methods that have been proposed in the last few decades by several authors.^{42,46,53–68} This lack of consensus on RCCS values causes the aforementioned ambiguity of SCSs. On the other hand, experimental aspects such as CS referencing, and signal assignment also contribute to the uncertainty of SCS values.^{69,70} Based on all this, the arising questions are: how much does the RCCS-related ambiguity influence secondary structure determination and what can be done to eliminate or at least reduce this effect? To address this problem, a comparative study of different RCCS datasets and calculation methods is necessary. Only very few works focus explicitly on the comparison of different RCCS prediction methods.^{71–75} Usually, such issues constitute marginal parts of the papers introducing new predictors.^{58,60,63–66,76}

We intend to fill this gap and we propose to discuss RCCS predictor development as a calibration problem. We give an overview of the theoretical and experimental background of the presently available RCCS predictors, focusing on their differences. Further on, we provide case studies demonstrating how the different RCCS prediction methods influence the secondary structure or structural propensity assessment of a protein. As examples, we chose well-known and extensively studied proteins: the folded ubiquitin, and α -synuclein and the trans-activation domain of p53 as IDPs, highlighting at the same time the effect of experimental conditions at various pH values and different temperatures. By means of statistical inference, we try to determine which RCCS predictor, if any, best represents the consensus of multiple predictors for a given experimental dataset. In the light of all these, one can choose and apply predictors simultaneously, a so far uncommon – but useful – practice.

2. Determining RCCSs: an ill-defined calibration problem

As purely computational approaches for determining RCCSs have been limited,^{77–81} producing an RCCS calculation method turns out to be a calibration process. Differences between RCCS prediction methods can be categorized as conceptual and



experimental. The conceptual properties of the presently available RCCS calculation methods are: the type of example system(s) chosen (*i.e.* small peptides, IDRs, IDPs), factors that are included in calculating RCCSs, such as local sequence, pH, temperature, ionic strength, the form of the equations providing RCCS values and the method used to parametrize these equations. The experimental differences arise primarily from the actual example systems chosen and to a lesser extent from the measurement uncertainty of chemical shifts. It has been pointed out that already the random coil state of polypeptides has to be clarified.⁸²

Historically, two main conceptual approaches were considered for the calibration of RCCSs. One approach is based on designing small peptides whose behavior is assumed to best represent the most disordered state any polypeptide might adopt.^{53–56,58,64,68,76} The other approach involves compiling a protein chemical shift database followed by a statistical analysis of the data.^{42,46,57,59–63,65–67} This approach has become increasingly popular with the growing number of IDP-related datasets in the Biological Magnetic Resonance Databank (BMRB).

Following the choice of suitable model systems, another issue is how to take experimental conditions into account. So far, according to the literature, temperature and pH, have been directly and ionic strength indirectly considered.^{62,65,66,76,83} Besides these experimental parameters, the local amino acid sequence has an impact on the CSs of an individual residue in the polypeptide chain and has been accounted for in some methods.

The small peptide approach has the advantage that the tabulation of RCCSs is very straightforward and requires no or very little computation. Also, one has extensive control over experimental conditions as the respective values of pH, temperature and ionic strength may all be precisely adjusted. On the other hand, it is much more difficult to cover the local compositional space of proteins. For example, if 20 amino acids and only the nearest neighbors are considered, 20^3 , meaning 800 combinations must be examined. Accounting for the neighboring ± 2 amino acids, this number jumps to 3.2 million. As it would be very time-consuming and costly to produce so many different peptides, in studies done so far, authors designed a given polypeptide frame (for example Gly–Gly–Xxx–Ala–Gly–Gly)⁵⁶ and varied a single amino acid in a central position. The sequential effect of the different amino acids on their neighboring partners is evaluated by their effect on the amino acids of the polypeptide frame, in the abovementioned case on glycines and alanine. On the other hand, the effect of a given amino acid on its neighbors depends also on the identity of the neighbors. Such pairwise and n-wise relationships are impossible to account for by the small peptide approaches utilized so far.

In contrast, database-related statistical approaches have the opposite strengths and weaknesses. With large numbers of CSs available for extensive numbers of proteins, the compositional space is much better covered than in peptide-based studies. The same local sequence may appear numerous times, therefore chemical shift values for all the involved amino acids are

observed numerous times as well. A large enough database even enables the determination of pairwise or n-wise correction terms for the effect of the local sequence. The drawback is, that the effect of experimental conditions is generally difficult to account for, as these parameters usually vary from entry to entry. Also, since database approaches directly use chemical shifts of proteins for calibration, it could be argued that the resulting RCCS values are more appropriate for studying proteins than RCCSs originating from small peptide studies. However, even if chemical shift data of IDPs are used,^{63,65,66} it is not guaranteed that all CS are RCCSs because of residual structural motifs in IDPs.⁸⁴ This requires authors to filter the data in some manner that decomposes the measured CSs into RCCSs and the different contributions of all the experimental conditions, the local sequence and, most importantly, residual structure.^{63,66,85,86} Loop regions, denatured proteins and even some peptides have been shown to not be completely disordered.^{87–102}

In the light of the above, in Table 1 we summarize the works that have been carried out with the aim of calibrating RCCSs. One can observe that database-derived, statistical approaches have recently been gaining popularity. It is interesting to note, that, except for the work of Bundi *et al.*,⁵⁴ systematic pH and temperature corrections only became available in the 2000's, while the effect of local sequence was already considered in the work of Braun *et al.*⁵⁵ in 1994. Sequence corrections became more elaborate in later RCCS predictors.

In this work, we focus on investigating free proteins in aqueous solutions and under the typical conditions of NMR studies. We are aware of works concerning RCCSs under high pressure,^{103–107} for phosphorylated,¹⁰⁸ posttranslationally modified¹⁰⁹ amino acids and in the presence of organic solvents,^{110,111} however we are not discussing these specialties here.

Below, we provide a brief overview of the approaches regarding RCCS corrections for the three aforementioned factors: local sequence, pH and temperature.

2.1. Sequence correction of RCCSs

Nearest neighbor-effects on the local structure and energetics of proteins is an extensively studied topic especially in the field of IDPs.¹¹²

The first sequence corrections of RCCSs were suggested by Braun *et al.*⁵⁵ Considering the Gly–Gly–Xxx–Ala and Gly–Gly–Gly–Ala sequences, corrections were defined as:

$$\Delta\delta^X = \delta_{A,N}^{GGXA} - \delta_{A,N}^{GGGA} \quad (2)$$

where X is any of the 20 naturally occurring amino acids, $\delta_{A,N}^{GGXA}$ and $\delta_{A,N}^{GGGA}$ are the chemical shift values of Ala in the corresponding peptides, $\Delta\delta^X$ is the sequence correction term for amino acid X. The N subscript refers to the amide nitrogen atom type. This definition approximates the effect of X on residue $i + 1$ by calculating this effect for alanine and setting the contribution of glycine as 0 ppm.

Although the calculation is very straightforward, this approach assumes that the local sequence effect experienced by alanine because of residues X is equal to what the remaining 19



Table 1 Features of available RCCS datasets and corresponding calculation methods. If the RCCS dataset/method had no specific name, the surname of the first author was used. The first letter of an author was included for differentiating two authors with identical surnames. Methods selected for our calculations are shown in bold

Method	Year	Ref.	Type of system	Corrections			Atom types					
				Sequence	Temperature	pH	H ^N	H ^α	C ^α	C ^β	C ^γ	N
McDonald	1969	53	Free amino acids, different small peptides	X	X	X	X	✓	X	X	X	X
Howarth	1978	67	Peptides and denatured proteins in D ₂ O	X	X	X	X	X	✓	✓	✓	X
Richarz	1978	68	Small peptide (GG-X-A)	X	X	X	X	X	✓	✓	✓	X
Bundi	1979	83	Small peptide (GG-X-A)	X	X	X	✓	✓	X	X	X	X
CSI	1992, 1994	41 and 42	Globular proteins	X	X	X	✓	✓	✓	✓	✓	✓
Braun	1994	55	Small peptide (GG-X-A)	✓	X	✓	X	X	X	X	X	✓
Wishart	1995	56	Small peptide (GG-X-A/P-GG)	✓	X	X	✓	✓	✓	✓	✓	✓
Lukin	1997	57	Database (BMRB and literature)	X	X	X	✓	✓	✓	✓	✓	✓
Schwarzinger	2000	58	Small peptides, in acidic 8 M urea	✓	X	X	✓	✓	✓	X	✓	✓
PSSI	2002	59	Database (selected BMRB)	X	X	X	✓	✓	✓	✓	✓	✓
Wang	2002	60	Database (selected BMRB)	✓	X	X	✓	✓	✓	✓	✓	✓
RefDB	2003	61	Selected BMRB data (database)	X	X	X	✓	✓	✓	✓	✓	✓
Wang L.	2006	46	Proteins from refDB	X	X	X	X	X	✓	✓	X	X
Camcoil	2009	62	Loop regions of globular proteins (selected BMRB)	✓	X	✓	✓	✓	✓	✓	✓	✓
ncIDP	2010	63	IDPs (mostly BMRB)	✓	X	X	✓	✓	✓	✓	✓	✓
Kjaergaard	2011	64 and 76	Small peptides	✓	✓	✓	✓	✓	✓	✓	✓	✓
Prosecco	2017	65	IDPs (selected BMRB)	✓	X	✓	✓	✓	✓	✓	✓	✓
Potenci	2018	66	IDPs (ncIDP extended)	✓	✓	✓	✓	✓	✓	✓	✓	✓

amino acids would experience, which is not necessarily the case. Also, taking only the preceding residue into account might be plausible for amide nitrogen, but not for other atom-types of the peptide backbone.⁶⁶

A similar approach was used in the work of Wishart *et al.*⁵⁶ considering Gly-Gly-Xxx-Ala-Gly-Gly and Gly-Gly-Xxx-Pro-Gly-Gly hexapeptides. Sequence correction was given as:

$$\Delta X = \delta X_A - \delta X_P \quad (3)$$

Here ΔX is the sequential correction for residue X , δX_A and δX_P are the chemical shift values of residue Xxx in Gly-Gly-Xxx-Ala-Gly-Gly and Gly-Gly-Xxx-Pro-Gly-Gly, respectively. In this case only the effect of proline on the preceding residue is considered. Although the Wishart dataset contains various individual correction terms, generally only atom-type wise averages of these are used in applications. This convenient practice obviously limits the accuracy of SCS calculation.^{38,113} Later small peptide-based works of Schwarzinger *et al.* and Kjaergaard *et al.* followed a similar experimental approach but used individual correction terms instead of averaging in calculations.^{58,64,76}

In 2000, Schwarzinger *et al.* used Gly-Gly-Xxx-Gly-Gly constructs and provided sequence correction terms for all four of the $i - 2$, $i - 1$, $i + 1$ and $i + 2$ positions.⁵⁸ These correction terms were respectively calculated as the CS difference of Gly1, Gly2, Gly4 and Gly5 in the Gly-Gly-Xxx-Gly-Gly and the reference Gly-Gly-Gly-Gly peptide according to the set of equations below

$$A = \delta(G1) - \delta(G1_{ref}) \quad (4)$$

$$B = \delta(G2) - \delta(G2_{ref}) \quad (5)$$

$$C = \delta(G3) - \delta(G3_{ref}) \quad (6)$$

$$D = \delta(G4) - \delta(G4_{ref}) \quad (7)$$

where $\delta(Gi)$ is the chemical shift of the Gly residue in position i in the Gly-Gly-Xxx-Gly-Gly peptide and $\delta(Gi_{ref})$ is the chemical shift of the same Gly residue in the reference Gly-Gly-Gly-Gly peptide. Each of the 20 amino acids has its set of A , B , C and D values. Thus, the sequence corrected RCCS of residue R can be calculated as follows.

$$\delta_R(\text{corrected}) = \delta_{\text{random}}(R) + A + B + C + D \quad (8)$$

Here, $\delta_{\text{random}}(R)$ is the uncorrected RCCS of residue R . This approach assumes that CSs of glycine are representative of all 20 amino acids in experiencing the sequential presence of residue Xxx . Despite using peptides with acetylated N-terminus and amidated C-terminus, the validity of $i - 2$ and $i + 2$ correction terms remains questionable because of so-called “end-effects”.^{114,115} Also, no sequence correction terms could be determined for the C^β atom-type because of the glycine frame used.

The first statistical approach in sequence correction was given by Wang *et al.*⁶⁰ They used more than 200 000 chemical shifts from BMRB to calibrate average random coil chemical shifts, pairwise sequence correction terms, average secondary structural chemical shift terms and terms for pairwise sequence effects in the random coil, β -strand and α -helical states. We



note, that in this case the definition of RCCS heavily depends on the identification of residues in the random coil state by means of VADAR, DSSP and PSSI.^{59,116,117} Using this definition of disorder, parametrization of eqn (9) and (10) could be performed:

$$\Delta(^X Y)_{n,s} = \langle \delta_{n,s}(X) \rangle - \langle \delta_{n,s}(w/o X) \rangle \quad (9)$$

$$\Delta(Y^Z)_{n,s} = \langle \delta_{n,s}(Z) \rangle - \langle \delta_{n,s}(w/o Z) \rangle \quad (10)$$

where $\Delta(^X Y)_{n,s}$ and $\Delta(Y^Z)_{n,s}$ are the contributions of preceding X and succeeding Z residues to the CS of atom-type n in residue Y, and in structural state s (helix, strand, random coil). $\langle \delta_{n,s}(w/o X) \rangle$ and $\langle \delta_{n,s}(w/o Z) \rangle$ are the corresponding average CS terms for cases where residue Y is not preceded by X and not followed by Z.

The next method including sequence correction terms in RCCS calculation was Camcoil,⁶² published by De Simone *et al.* in 2009. Therein, a database of 1772 BMRB entries belonging to proteins possessing PDB entries was used. Residue specific RCCSs were calculated as the average CSs of the given residue found in the coil regions of the considered proteins. Pairwise correction contributions for the preceding and succeeding residue were calculated by averaging for CSs of the appropriate residue pairs in the database. Moreover, a set of atom-type dependent weight factors for these correction terms was proposed for the predicted RCCS of atom *i* of residue A in a BAC peptide triplet:

$$\delta_{iA}^{RC} = \delta_{iA}^0 + \alpha_i^- \delta_{iBA}^1 + \alpha_i^+ \delta_{iAC}^1 \quad (11)$$

Here δ_{iA}^0 is the uncorrected RCCS of atom *i* of residue A, calculated as the database average. Similarly, δ_{iBA}^1 , δ_{iAC}^1 are the sequence correction terms of residues B and C calculated as averages, while α_i^- and α_i^+ are corresponding weight factors for atom-type *i* and for the preceding and succeeding positions. Weight factors were optimized by minimizing the deviation of predicted CSs from the experimentally determined CSs of a set of proteins under denaturing conditions. In practice, Camcoil is available as an online application.¹¹⁸

The ncIDP method by Tamiola *et al.*, includes sequence corrections.⁶³ This is achieved by solving a set of equations of the following form:

$$\delta^n(x,a,y,i) = \delta_{RC}^n(a) + \Delta_{-1}^n(x) + \Delta_{+1}^n(y) + \varepsilon^n(i) \quad (12)$$

using singular value decomposition.

Here, the tripeptide *xay* is considered, and $\delta^n(x,a,y,i)$ is the CS of atom-type *n* of the *i*-th residue, $\delta_{RC}^n(a)$ is the uncorrected RCCS of *a*, $\Delta_{-1}^n(x)$ and $\Delta_{+1}^n(y)$ are sequence correction terms for the preceding and succeeding residue, while the $\varepsilon^n(i)$ term accounts for any deviation caused by pH, temperature or CS referencing of individual datasets. Thus, the resulting sequence correction terms – despite originating from a database approach – are not pairwise and depend only on the identity of the preceding and succeeding residue. In practice, ncIDP is available online.¹¹³

In 2011, Kjaergaard *et al.* provided an RCCS dataset including sequence correction terms derived from the investigation of Gln–Gln–Xxx–Gln–Gln peptides.⁶⁴ The procedure of Schwarzinger *et al.* was adopted, however Gln–Gln–Gln–Gln–Gln was defined as the reference peptide. Correction terms for sequence positions *i* – 2, *i* – 1, *i* + 1 and *i* + 2 and for the H^N, H^α, N, C^α, C^γ and C^β atom types were determined. RCCS calculation follows eqn (4)–(8) with the adjustment of the reference peptide. Problems mentioned earlier, regarding the validity of correction terms for the two terminal positions, are present here as well. This RCCS calculation method is also available as a web application.¹¹⁹

In the first attempt to apply an advanced machine learning approach for RCCS calibration, Sanz-Hernández and De Simone built the Prosecco neural network model, which uses sequence correction terms.⁶⁵ A sufficiently large dataset of more than 20 000 CSs of IDPs and IDRs from BMRB was used, and determination of pairwise correction terms for the *i* – 2, *i* – 1, *i* + 1 and *i* + 2 positions was achieved. The calculation involved the use of smoothed empirical probability density functions of CSs derived by applying Gaussian kernels according to eqn (13).

$$\hat{\delta}_{ij}^A(\delta) = \frac{1}{n_{ij}^A} \sum_{l=1}^{n_{ij}^A} G_K(\delta - \delta_l) \quad (13)$$

where $G_K(\delta - \delta_l)$ is a Gaussian kernel function centered at the experimental chemical shift δ_l ; n_{ij}^A is the number of cases when residue *j* is found in the given relative position with respect to residue *i*, while $\hat{\delta}_{ij}^A(\delta)$ is the resulting smoothed empirical probability density function of pairwise chemical shifts. From these empirical probability density functions, the corresponding $\Delta\delta_{ij}^A$ sequence correction term is calculated as:

$$\Delta\delta_{i,j}^A = \hat{\delta}_{i,j}^A - \delta_i^A \quad (14)$$

where $\hat{\delta}_{i,j}^A$ is the expected value of the $\hat{\delta}_{ij}^A(\delta)$ empirical probability density function, δ_i^A is the uncorrected RCCS of residue *i*, which is defined, similarly, as the expected value of the smoothed empirical probability density function of the chemical shifts of residue *i* in the database.

The correction terms of Prosecco are not used directly but are multiplied by weight factors $w_{i,k-2}^A$, $w_{i,j-1}^A$, $w_{i,l+1}^A$, $w_{i,m+2}^A$ defined as the corresponding empirical negative overlap integral of the corresponding primary and pairwise probability density functions of the CSs of the central amino acid in a quintuple of residues:

$$c_{seq} = \frac{1}{N_W} \left(w_{i,k-2}^A \Delta\delta_{i,k-2}^A + w_{i,j-1}^A \Delta\delta_{i,j-1}^A + w_{i,l+1}^A \Delta\delta_{i,l+1}^A + w_{i,m+2}^A \Delta\delta_{i,m+2}^A \right) \quad (15)$$

where c_{seq} indicates the complete sequence correction term of the RCCS, N_W is a general weight factor which is intended to scale the respective contributions of the sequence correction terms and of the uncorrected RCCSs. Its value was determined in the optimization procedure yielding the Prosecco model. The exact parameters of the neural network model have not been



published, and Prosecco can only be used as a web application.¹²⁰

Presently the latest RCCS prediction method with sequence correction terms is Potenci by Nielsen and Mulder, published in 2018.⁶⁶ The local sequence effect is accounted for from position $i - 2$ to position $i + 2$. The formulation of sequence correction terms combines an earlier idea with a novel one. As seen in both small peptide studies and the practical applications of the Wang method, general correction terms could theoretically be assigned to all 20 amino acids for all atom-types. Nielsen and Mulder determined a set of such general correction terms for all 6 canonical atom-types, and H β . Their novel idea was that the meticulous determination of pairwise correction terms can be neglected, instead, so-called correlated amino acid or second order contributions are determined. To achieve this, the 20 amino acids were divided into 7 groups according to the properties of the side-chain: G Gly, P Pro, r aromatics (Phe, Tyr, Trp), a aliphatics (Leu, Ile, Val, Met, Cys) and A, "+" positive (Lys, Arg), "-" negatives (Asp, Glu), while p polar residues (Asn, Gln, Ser, Thr, His). The direct neighbor correction and next-neighbor correction terms were defined using a principal component representation of amino acids suggested by Georgiev¹²¹ and corresponding w_j^k tunable weights:

$$\Delta(p) = \sum_{k=-2,-1,1,2} \Delta_k(a_{i+k}) \quad (16)$$

$$\Delta_k(a_{i+k}) = \sum_{j=1}^{\gamma_k} w_j^k a_j^k(a_{i+k}) \quad (17)$$

where $\Delta(p)$ is the general neighbor correction built up from the $\Delta_k(a_{i+k})$ individual contributions of the neighboring amino acids in all four of the $i \pm 2$ positions. These individual terms comprise linear combinations of the $a_j^k(a_{i+k})$ principal component values of the first γ_k Georgiev principal components, after multiplication by the tunable w_j^k weights. During the fitting procedure, the γ_k number of principal components to be used and the values of w_j^k were optimized resulting in the ultimate Potenci model. The C- and N-terminal residues were treated as two separate residue-types. Optimizing γ_k , enabled the use of a smaller number of adjustable parameters than a theoretically extensive model would have to include.

In determining correlated contribution terms, a similar approach was followed. For each atom-type, $\omega_{i,m}^k$ was defined as the contribution of residue type m to the CS of residue type l in position i when m is in relative position k with respect to i :

$$\chi(p) = \sum_{k=-2,-1,1,2} \chi_k(a_i, a_{i+k}) \quad (18)$$

$$\chi_k(a_i, a_{i+k}) = \begin{cases} \omega_{g(a_i),g(a_{i+k})}^k & \text{if } (k, g(a_i), g(a_{i+k})) \in \Pi \\ 0 & \text{else} \end{cases} \quad (19)$$

Here $\chi(p)$ is the complete correlated residue contribution to the predicted RCCS. The complete contribution is built up from the $\chi_k(a_i, a_{i+k})$ individual contributions, each of which are built up from the adjustable $\omega_{g(a_i),g(a_{i+k})}^k$ parameters if the given correlated contribution is significant, otherwise they are zero.

Accordingly, Π denotes a certain set of values for k , $g(a_i)$ and $g(a_{i+k})$. For any atom-type, theoretically $4 \times 7 \times 7$ such parameters exist, as there are 4 relative positions ($i - 2, i - 1, i + 1, i + 2$) and seven groups of amino acids. In the optimization process, the number of such terms to be used was also treated as an adjustable parameter, and its ultimate value was smaller than the theoretical maximum for each atom type.

2.2. pH correction of RCCSs

The importance of pH was already noted in the earliest efforts for RCCS determination. Despite this, only a few RCCS predictors apply pH correction, even less treat the pH as a quasi-continuous variable. Generally, the non-structure related effect of pH on CSs is only important in the case of residues with titratable sidechains: glutamic acid, aspartic acid, histidine and their direct neighbors are affected in the acidic and neutral regime.^{122,123}

In 3 early works Richarz and Wüthrich,⁶⁸ Bundi and Wüthrich⁵⁴ and Braun *et al.*⁵⁵ respectively published ¹H, ¹³C and ¹⁵N chemical shifts of Gly-Gly-Xxx-Ala tetrapeptides. These three works established the first so-called binary pH-correction scheme for RCCSs. That is, respective pairs of RCCSs were proposed for Glu, Asp and His residues: one for pH values at least 1.5 smaller than the pK_a of the given residue, and one for pH values at least 1.5 larger than the corresponding pK_a . This is equivalent to providing an RCCS value for the completely protonated and the completely deprotonated state of the residue, respectively. For Asp and Glu residues, this might be a smaller issue as peptide and protein studies are either carried out under very acidic condition (pH < 3), where even the acidic sidechains of these two amino acids are almost completely protonated, or, at neutral pHs, where both are close to completely deprotonated.¹²²⁻¹²⁵ In contrast, histidine has typical pK_a values between 6 and 7 and is therefore partially protonated under close to physiological conditions, making it fall into a "blind range" of binary pH-correction schemes.

Camcoil, published in 2009 (ref. 62) also has a binary pH-correction scheme. That is, spectra recorded under either acidic (pH < 2) or close to neutral (average pH = 6.1) conditions were used. The correction terms for acidic conditions, nominally pH = 2, were optimized by fitting chemical shifts of aspartic acid, glutamic acid and histidine residues of two BMRB entries acquired under denaturing conditions at acidic pH values.

In 2011, Kjaergaard *et al.* took a more sophisticated approach to performing pH-correction of RCCSs⁷⁶ using Gly-Gly-Xxx-Gly-Gly pentapeptides similarly to Schwarzingler *et al.*⁵⁸ In contrast to the original work, Kjaergaard *et al.* determined RCCS values at pH = 6.5 instead of pH = 2.5 in 8 M urea. Moreover, they performed pH-titration of the peptides with Xxx = Asp, Glu and His and determined the side chain pK_a values of these residues by non-linear fitting of the titration curves. The equation for the $\delta(\text{pH})$ pH-corrected RCCS is the linear combination shown by eqn (20).

$$\delta(\text{pH}) = \delta_A \frac{K_a}{10^{-\text{pH}} + K_a} + \delta_{\text{HA}} \left(1 - \frac{K_a}{10^{-\text{pH}} + K_a} \right) \quad (20)$$



Here K_a is the acid dissociation constant of the side chain, δ_A and δ_{HA} are the RCCS values corresponding to the completely deprotonated and completely protonated state, respectively.

On the basis of eqn (20), originating from the classical equation for chemical equilibria, a continuous pH-correction is enabled.

Such an approach is dependent on the knowledge of side chain pK_a values for individual residues in the protein.

Another limitation is that in practice, the pH-dependence of the chemical shifts of titratable residues in proteins often follows a Hill-model,^{122,124,126} including an extra parameter called the Hill-coefficient. The advantage is the lack of a “blind range” as opposed to the earlier, binary corrections.

The Prosecco server provides a binary pH-correction with the same parametrization of pH correction terms as Camcoil.^{62,65} In Prosecco, 6 BMRB entries were used to optimize the correction terms, with chemical shifts corresponding to nominal pH values of 2.8 and 6.4. During optimization the minimization of the absolute difference between calculated and experimentally determined chemical shifts of the 6 analyzed IDP entries was performed.

Being a very synthetic approach, the pH-correction of Potenci utilizes earlier ideas, but introduces some novelty to the field.⁶⁶ Potenci's uncorrected RCCSs correspond to pH = 7.0. Therefore, pH-correction is only needed if the pH differs from this value. The correction considers a linear combination approach similar to eqn (20). The novelty is, that the Hill model is utilized, and the f^{HA} relative concentration of the protonated side chain is calculated as in eqn (21).

$$f^{HA} = \frac{10^{n_H(pK_a - pH)}}{1 + 10^{n_H(pK_a - pH)}} \quad (21)$$

Here n stands for the Hill-coefficient.

Further on, using the determined relative concentrations, the final pH-correction is acquired according to eqn (22).

$$\varepsilon_k(a_{i+k}, pK_{a_{i+k}}, pH) = \Delta\delta_{HA-A}^k(a_{i+k})(f^{HA}(pH) - f^{HA}(pH = 7)) \quad (22)$$

Here $\varepsilon_k(a_{i+k}, pK_{a_{i+k}}, pH)$ is the pH correction applied for amino acid a in the $i + k$ relative position with respect to a residue with titratable side chain and nominal acid dissociation constant $K_{a_{i+k}}$ at a given pH. $\Delta\delta_{HA-A}^k$ is the chemical shift difference between the completely protonated state and the completely deprotonated state of the titratable side chain. The main novelty is that the effect of the protonation state of a titratable residue on its nearest neighbors is accounted for. The $\Delta\delta_{HA-A}^k$ values are taken from the work of Platzer *et al.*¹²³ Both Hill-coefficients and the specific pK_a values of titratable residues in the given sequence are calculated by pepKalc.¹²⁶ Since pepKalc requires the ionic strength of the sample, ionic strength is an additional input of Potenci.

2.3. Temperature correction of RCCSs

Although the effect of temperature on chemical shifts is well-known and widely studied,¹²⁷⁻¹³⁰ at present, there are only two RCCS calculation methods that explicitly take this effect into account by introducing correction terms. In the case of small

peptide derived RCCS datasets, the temperature of the experiments is known precisely, so the RCCS values correspond to this temperature, however, it is not straightforward, how one should use such data for measurements that were carried out at different temperatures. On the contrary, for database-derived predictors the opposite is true. Although these works usually use measurement data covering a relatively wide range of temperature values, it is not clear, what temperature the acquired RCCSs correspond to. Usually, the average temperature of the different spectra is reported however, this still does not solve the problem of extrapolation.

The first continuous temperature-correction terms in RCCS calculation were introduced by Kjaergaard *et al.*⁷⁶ The chemical shifts of Gly-Gly-Xxx-Gly-Gly peptides were determined at 5, 15, 25, 35 and 45 °C. As in all cases, a linear dependence was observed, the slopes of the corresponding curves were obtained from linear fitting. Thus, temperature coefficients for the H^N , H^α , N , C^α , C' and C^β atom types of the 20 amino acids are available and the temperature correction is performed as:

$$d_{T_{corr}} = a \times (T - T_{ref}) \quad (23)$$

where T_{ref} is the reference temperature of the uncorrected RCCS, a is the temperature coefficient and $d_{T_{corr}}$ is the correction term to be applied to obtain the corrected RCCS at temperature T . Note, that the temperature coefficients calibrated on short peptides might only be best guesses for proteins.

Once the reference temperature of any RCCS dataset is available, it is theoretically possible to transfer temperature coefficients from this RCCS prediction method to others - in stark contrast to sequence correction terms. Nielsen and Mulder performed such a transfer of temperature coefficients published by Kjaergaard *et al.* when developing Potenci. Accordingly, Potenci uses eqn (23) with $T_{ref} = 298$ K for temperature correction.

3. Importance and non-equivalence of RCCS predictors in practical applications

We proposed to investigate how the selection of a given RCCS predictor influences the identification of secondary structural motifs and structural propensities. To test the performance of RCCS predictors we chose three well-studied proteins: the folded ubiquitin and two IDPs: α -synuclein and a 60-residue part of the transactivation domain (TAD) of p53. For ubiquitin and α -synuclein chemical shift data are available from the BMRB under codes 4769, 27348 and 18857, and these systems were tested in our lab, too. The p53TAD¹⁻⁶⁰ construct has been studied in detail by our group earlier.^{131,132} In the followings, from the available atom types, we limit ourselves to C^α environments. Chemical shifts of C^α are generally considered one of, if not the, most sensitive parameters to secondary structure and structural propensities and have been preferred over their counterparts in various cases.^{3,37,46,52,59,133-140} Out of the numerous RCCS predictors in Table 1, we selected those 8



which have at least some sort of local sequence correction, yield prediction for the C^α atom type and could theoretically be considered appropriate for studying IDPs in aqueous solutions.

Comparison of RCCS predictors was performed in two different ways. First, by visual analysis of the calculated C^α SCS plots where one has to focus on regions showing meaningful differences depending on the RCCS predictor, and therefore leading to the controversial identification of secondary structural propensities. Second, *via* comparison of the predictors by a versatile non-parametric statistical tool: the sum of ranking differences and comparison of ranks to random numbers (SRD-CRRN) method augmented by one-way analysis of variance (ANOVA) with *post hoc* Bonferroni pair-wise tests.^{141–144}

3.1. The visual comparison of RCCS predictors for C^α chemical shifts

3.1.1. A folded protein example: ubiquitin. To test different RCCS predictors on a folded protein, we chose the well-known 76-residues long ubiquitin. Human and yeast ubiquitin have similar structures. Yeast ubiquitin has P19S, E24D and A28S mutations but these do not influence the overall structure. The 3D structural models determined by different methods are illustrated in Fig. 1. As observed, the structures are similar, differences concern mainly the limits of certain structural motifs – proving that already different 3D structure elucidation approaches do not give completely identical results.

Using the chemical shift information, the secondary structural elements can be assessed, and for this purpose we prepared the C^α SCS plots of ubiquitin (BMRB: 4769) with the selected 8 predictors (see Fig. 2). A simple visual inspection shows that there is little discrepancy between the different methods, as also represented in the plot showing the median values. Regions with a given secondary structure have high (± 2 ppm) C^α SCS values, while the mobile loop regions show ± 0.5 ppm values, that are characteristic of the behavior of IDPs. We use the median instead of the mean to represent the general

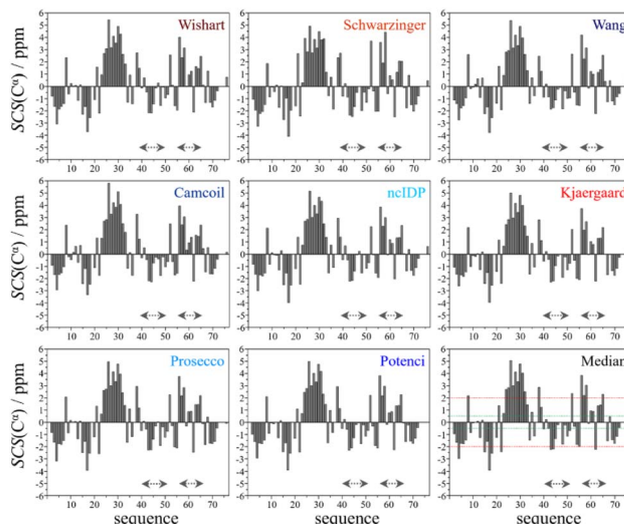


Fig. 2 C^α SCS plots of ubiquitin, based on data from BMRB 4769; pH = 7.5, $T = 303$ K – for the chosen RCCS predictors and taking the median. Small-peptide RCCS predictors (Wishart, Schwarzingler, Kjaergaard) are shown in shades of red, while database-derived ones (Wang, Camcoil, nCIDP, Prosecco, Potenci) are shown in shades of blue.

trend of the 8 selected RCCS predictors because the former is less sensitive to outliers. As will be shown later, outliers do appear when SCS values with different RCCS predictors are calculated, especially in the case of IDPs.

Still, despite the general similarity of plots in Fig. 2, some differences can be noticed. The Gln40–Glu51 region, which is highlighted on the sequence in Fig. 1, is suggested by structure elucidation to host two β -structures separated by a short loop. On the SCS plots, this structural motif is supported by most methods, appearing as an “inverse valley”, but this pattern is visually less well-defined for the Camcoil predictor. The discrepancy originates from the SCS values of Gly47 being either

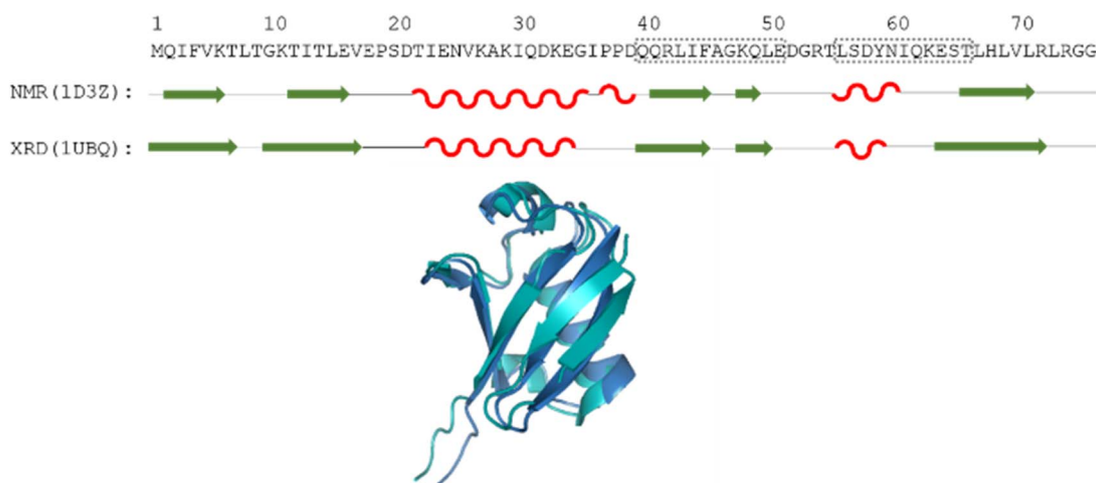


Fig. 1 The amino acid sequence of human ubiquitin; structures determined by different methods, where secondary structural elements are represented by red wave (helix), green arrow (sheet) and black line (flexible loop) below the sequence; as well as the aligned 3D structures (PDB: 1D3Z, cyan, PDB: 1UBQ, dark blue). Regions Gln40–Glu51 and Leu56–Thr66 are highlighted by dotted grey boxes.



sophistication, may rely on. While all the plots in Fig. 4 differ from each other to some extent, C^α SCS plots yielded by the Schwarzingger and Camcoil methods are especially unique. In the case of the Schwarzingger method, the spikes appearing at Glu and Asp residues result from the corresponding RCCS values having been recorded at pH = 2.3. The titratable side chains of these residues were completely or close to completely protonated under the circumstances of RCCS calibration, while the same side chains were completely deprotonated at pH = 6.5, where the data of BMRB entry 27 348 were recorded. In the case of Camcoil, the surprising feature is that the average amplitude of the SCSs is generally larger than for the other methods, irrespective of residue type. We see no direct connection between this and the theoretical background of Camcoil. However, one must be aware of this feature when using SCSs calculated by Camcoil either directly or by deriving structural probabilities from them in the $\delta 2D^{35,148}$ method.

For the other 6 C^α SCS plots of Fig. 4, a few general tendencies can be noticed. A set of predominantly negative values at the C-terminal of the protein are seen in the plot corresponding to the Kjaergaard method with a similar pattern being present in the Wang plot, the Prosecco plot and, to a smaller degree, in the Potenci plot, too. The C^α SCS plot made using the Wishart data set indicates some sets of consecutive negative values in this sequential region, but these are separated by short sets of consecutive positive values. The positive spike appearing in the Wishart plot belongs to His50 and can be attributed to the effect of pH. The Wishart data set was collected at pH = 5.0, where titratable side chain of histidine was close to completely protonated, while at pH = 6.5 the same side chain is already partially deprotonated, resulting in an inherent and uncorrected error of SCSs.

A part of the sequence, where a structural propensity could be assumed to be present, is the Ala18–Gly36 region according to Wishart and between Ala19–Glu28 according to the Kjaergaard method. Data obtained from the Potenci method could also be argued to mildly reinforce this tendency.

The C^α SCS plot based on the Wang method is peculiar even in itself. Up until residue Gln99, positive SCS values, many of which exceed 0.4 ppm, dominate the plot, with very few negative values of smaller than 0.2 ppm amplitude breaking the trend. From Gln99 onward, negative values, indicating β -sheet propensity dominate. Because of regular breaks in the trends and considerable variation in the amplitude of consecutive SCSs of the same sign, the Wang plot of α -synuclein is a typical example of a C^α SCS plot, which is very difficult to interpret, even qualitatively.

On the contrary, the C^α SCS plot of α -synuclein by ncIDP has amplitudes below 0.3 ppm almost exclusively, and no longer

series of consecutive SCSs with the same sign are present. The plot is very similar to small amplitude white noise, as one would expect for a completely unstructured protein.

3.1.3. An IDP with pronounced structural propensity: the p53TAD¹⁻⁶⁰. To demonstrate differences between RCCS predictors we show an example of a disordered protein fragment p53TAD¹⁻⁶⁰ (Fig. 5) – studied extensively in our lab, and in the literature – where almost all predictors agree on the presence of residual structure but the location and type of these motifs is controversial.^{131,132}

In the median C^α SCS plot of Fig. 6, the Gln16–Lys24 region of p53TAD¹⁻⁶⁰, has a set of consecutive positive values, suggesting a helical propensity. However, both the strength and exact localization of this propensity vary between the individual predictors. The strongest suggestion for residual helicity originates from Camcoil and the Wishart method, but even these two plots differ considerably in their patterns. With the Schwarzingger method there are only two positive spikes at Asp21 and Leu22, and no propensity can be supposed. The double spikes appearing at Asp41–Asp42 and Asp48–Asp49 because of the acidic calibration of the Schwarzingger method should also be noticed. In the Val31–Asp40 region neither glutamate nor aspartate residues are found, yet the Schwarzingger plot suggests a relatively convincing β -propensity. Neither of the remaining methods reinforces this tendency in comparable strength and length. Some β -motif could also be

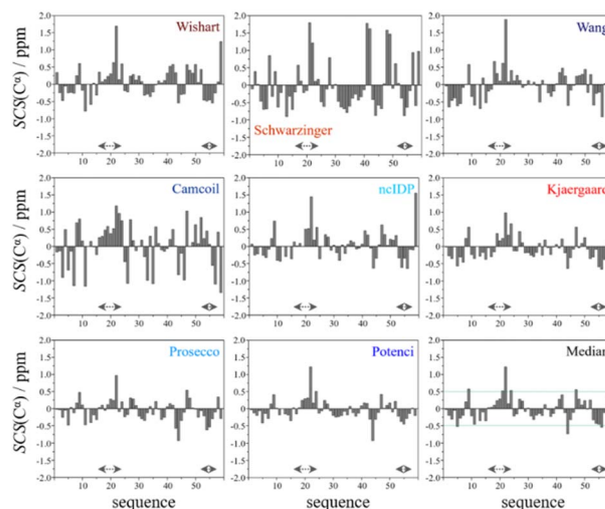


Fig. 6 C^α SCS plots of p53TAD¹⁻⁶⁰, data acquired in-house, pH = 6.0, $T = 313$ K. Small-peptide RCCS predictors (Wishart, Schwarzingger, Kjaergaard) are shown in shades of red, while database-derived ones (Wang, Camcoil, ncIDP, Prosecco, Potenci) are shown in shades of blue.



Fig. 5 Amino acid sequence of p53TAD¹⁻⁶⁰. The TAD1 (green) and TAD2 (light blue) regions have been color-coded. The N-terminal Gly and Ser residues denoted by smaller case letters were part of the construct but are not part of natural p53. Regions having been suggested to harbor helical propensities are highlighted by grey frames.



present close to the C-terminus between Trp53 and Asp57, according to the median plot, however the patterns of Camcoil and the Schwarzsinger method differ from those of their peers considerably in this aspect.

Summarizing the visual inspection for all the above-mentioned biomolecules, in the case of folded ubiquitin differences between RCCS predictors are not crucial. Also, assessing α -synuclein and p53TAD¹⁻⁶⁰ to be fundamentally unstructured is possible using each of the eight RCCS predictors. However, assessment of the presence or absence of regions with secondary structural propensities – the primary aim of SCS analysis – is not at all clear. The eight selected methods differ in both the amplitudes of SCS values, and in the corresponding trends of the signs thereof. Thus, evaluation of secondary structural propensities according to this set of RCCS predictors is ambiguous, raising various questions. Which RCCS calculation method(s) is one supposed to use for a given experimental dataset? If each method has its own limitations, is there a way to use a set of different RCCS predictors to get a realistic idea about these mild propensities? Is there a single RCCS prediction method which generally best represents the consensus of all 8?

3.2. The statistical comparison of RCCS predictors for C^z chemical shifts

Above, we have shown examples for differences between RCCS predictors which might completely alter the visual identification of residual structure in IDPs. Below, we show that some differences are statistically significant, and we highlight how statistics can be used to find predictors best representing the consensus of a predictor ensemble.

3.2.1. Statistical methodology. One conclusion of the qualitative picture is that – especially in the case of IDPs – the RCCS predictors are non-equivalent. Therefore, the question arises, whether the detected differences are significant, or not? To decide this, we chose the SRD-CRRN method – which is based on comparing a set of vectors to a reference vector – as this approach can give answers to our queries. To apply SRD-CRRN for the present protein studies, the SCS pattern for a single atom type (presently C^z) should be represented by a vector. Thus, the ensemble of the selected eight RCCS predictors will provide eight vectors. The so-called reference vector – to which these individual vectors are compared – is chosen to be the median vector of the corresponding C^z SCS dataset (represented in the median plots of Fig. 2, 4 and 6). This way SRD-CRRN shows how well individual RCCS predictors reproduce the consensus C^z SCS pattern of the whole ensemble. For SRD-CRRN calculation, the software is available at,¹⁴⁹ which will also generate plots automatically. A detailed and picturesque description of the SRD procedure can be found in ref. 143.

We illustrate the use of our approach on a model example, where five datasets were generated with thirty observations each. Note here, that for the SCS data, the only data pretreatment needed is the removal of missing SCS values for the missing assignments. The input data matrix contains the

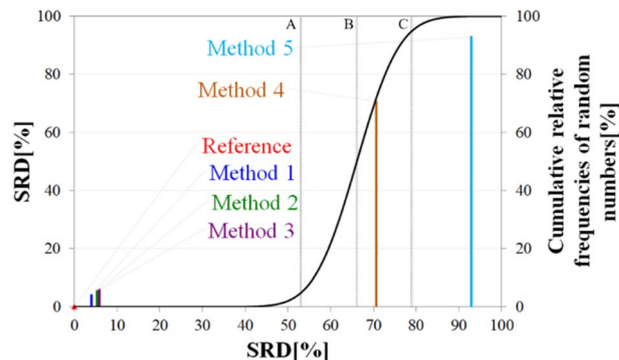


Fig. 7 General representation of SRD-CRRN results for a model example.

vectors corresponding to the methods to be compared. In the next step the SRD algorithm performs the comparison and validation by two built-in approaches. The first one is a comparison of SRD values to the SRD distribution of random vectors; and the second one is a 5-fold cross-validation. Running such an SRD-CRRN implementation results in a graph similar to the one in Fig. 7.

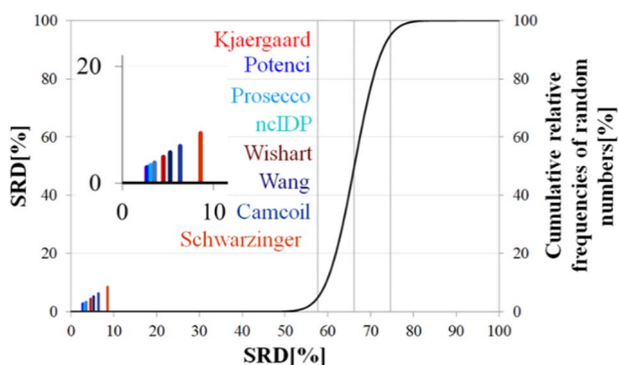
The theoretical cumulative distribution function of random vectors is represented by the black curve, and its numerical values are shown on the y-axis on the right. The curve is sigmoidal, as the SRD values of random vectors may be approximated by a normal distribution already for relatively small sample sizes (in our case thirty), as proved by Héberger and Kollár-Hunek.¹⁴² Lines A, B and C represent the 5th percentile, median and 95th percentile of this distribution, meaning the interval between A and C is the region of insignificance (insignificant region at $\alpha = 5\%$ level of significance). The compared five methods from our example are represented by colorful sticks. The height of a given stick is equal to its x-coordinate, that is the SRD score of the corresponding method (*i.e.* 4.24 for method 1). This is the reason for the SRD score being shown on both the x and the left y-axis. In the chosen model example, the reference method also appears in the SRD-CRRN plot. Obviously, the SRD value of this is 0 by definition, as its ranking is identical to that of the reference vector, meaning itself. Generally, in the SRD-CRRN applications this is not shown. Further on, Fig. 7 shows, methods 1, 2 and 3 reproduce the reference ranking much better than random numbers, as the corresponding colorful sticks are close to zero and they are far away from the region of insignificance A–C. In contrast, method 4 falls in the insignificant region, meaning it is not linked to the reference vector by any deterministic relationship. Method 5 is related to the reference vector, but produces a ranking inversely correlated to the reference, and as a result it is situated closer to the value of 100 than to the median of the distribution of random numbers. In conclusion, Fig. 7 provides an ordering of the compared methods according to their performance. Still, one can observe that methods 1, 2 and 3 are close to each other but there is no indication whether one is significantly better than the other. This can be decided by applying ANOVA on the data provided by the built-in 5-fold



Table 2 The one-way ANOVA and Bonferroni test result table for the model SRD-CRRN example

<i>p</i> (ANOVA)	Method	Mean SRD score	G1	G2	G3	G4
$<10^{-7}$	Method 1	4.24	****			
	Method 2	5.56	****	****		
	Method 3	6.14		****		
	Method 4	70.21			****	
	Method 5	92.78				****

cross-validation of the SRD algorithm. As $p(\text{ANOVA})$ in Table 2 is, $<10^{-7}$, which is $<5\%$, the test is significant, meaning not all methods perform equivalently well. This is highlighted by the mean SRD score of the methods shown in Table 2. These mean SRD score values correspond to the positions of the stick in Fig. 7. Which of the five methods are significantly better or worse can be checked by a Bonferroni *post hoc* test.¹⁴⁴ The Bonferroni test provides a grouping pattern of the methods as shown in Table 2. In this example, the chosen five methods form four homogenous groups named G1, G2, G3 and G4. The grouping pattern highlights that method 1 and 2 are not differentiable at a 5% level of significance by the *post hoc* Bonferroni *t*-test. Similarly, method 2 and 3 can be treated as equivalent, however method 1 performs clearly better than Method 3 based on the Bonferroni test. Method 4 and 5 are significantly different from all other methods, therefore they form independent groups G4, G5. This model example was

**Fig. 8** SRD-CRRN results for C^α SCS vectors of ubiquitin (BMRB 4769). Small-peptide RCCS predictors (Wishart, Schwarzsinger, Kjaergaard) are shown in shades of red, while database-derived ones (Wang, Camcoil, nCIDP, Prosecco, Potenci) are shown in shades of blue.

deliberately made to show that a method might belong to multiple homogenous groups as method 2 belongs to both groups G1 and G2; but it is also possible to have groups containing a single method like G3 and G4.

3.2.2. SRD-CRRN and ANOVA analysis of ubiquitin. For ubiquitin, the calculated SRD-CRRN plot is shown in Fig. 8. A simple inspection on the dispersion of sticks tells that the general consensus of the ensemble of predictors is strong, all the different methods are close to zero – as expected based on the high similarities concluded from the qualitative analysis of Fig. 2 plots.

Still, the RCCS predictors align in a certain order and according to ANOVA they are, even if seemingly not very different, not all equivalent. The Bonferroni *post hoc* test indicates that the two best methods, Potenci and that of Kjaergaard are equivalent at a 5% level of significance (Table 3). All other methods are significantly different from each other. The Schwarzsinger method falling visibly behind all the other predictors is explained by its calibration under acidic circumstances. This makes Schwarzsinger SCS values for residues with titratable side chains like Asp21, Asp24 and Asp32 much larger than those given by the other predictors. This effect might go unnoticed in visual assessment of Fig. 2 but is highlighted by SRD-CRRN.

Another interesting observation is, that four more recent methods – three database-derived (Potenci, Prosecco, nCIDP) and one small peptide-based (Kjaergaard) perform best, and they are followed by the small peptide-based method of Wishart. Thus, the Wishart method, which is the oldest one here, ends up in front of three methods developed later (Camcoil, Wang, Schwarzsinger) that could naively be considered improvements in the field.

3.2.3. SRD-CRRN and ANOVA analysis of α -synuclein. As could be expected based on the qualitative examination of C^α SCS plots, SRD-CRRN results for α -synuclein differ significantly from the picture obtained for ubiquitin (Fig. 9). The entire group of predictors is generally further away from zero, indicating weaker consensus regarding secondary structure. The much larger average of the entire set of SRD values for α -synuclein indicates that no method reproduces the consensus to such a degree as even the worst one in the case of the folded ubiquitin. This is exactly due to the intrinsically disordered nature of α -synuclein bearing weak structural propensities in its free form. Thus, small differences in RCCS prediction become important and their effect is reflected in the SRD analysis.

Table 3 The ANOVA and Bonferroni test results for SRD-CRRN data of ubiquitin, BMRB 4769

<i>p</i> (ANOVA)	Method	Mean SRD score	G1	G2	G3	G4	G5	G6	G7
$<10^{-7}$	Potenci	2.71	****						
	Kjaergaard	2.75	****						
	Prosecco	3.14		****					
	nCIDP	3.53			****				
	Wishart	4.61				****			
	Wang	5.31					****		
	Camcoil	6.50						****	
	Schwarzsinger	8.71							****



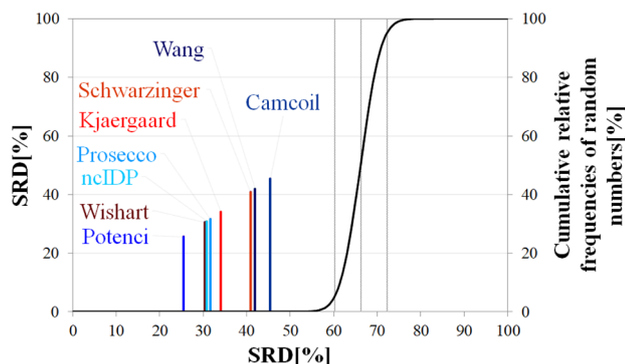


Fig. 9 SRD-CRRN results for C^α SCS vectors of α -synuclein (BMRB 27348). Small-peptide RCCS predictors (Wishart, Schwarzsinger, Kjaergaard) are shown in shades of red, while database-derived ones (Wang, Camcoil, ncIDP, Prosecco, Potenci) are shown in shades of blue.

A strongly phrased interpretation is that in the case of α -synuclein, although it is possible to find RCCS predictors which better reproduce the consensus of the eight considered methods, no single prediction is even close to being equivalent to this consensus SCS vector.

Differences in median SRD are larger, indicating a more pronounced ordering and grouping of the predictors. This is reinforced by the ANOVA *post hoc* analysis results (Table 4). Potenci performs best, finishing at first place, followed by the Wishart method, ncIDP and Prosecco. As for these three, at a 5% level of significance ncIDP – located in between – is indistinguishable from both the Wishart and Prosecco methods. In turn, these latter two are not equivalent according to the Bonferroni test. The remaining four methods form no homogenous groups but are all pairwise distinguishable from each other. This result highlights the usefulness of conducting ANOVA on the SRD data, as by a simple visual inspection of Fig. 9 one would consider the Wang and Schwarzsinger methods equivalent. The ordering of the methods differs from that found for ubiquitin; however, one has to observe that the first five and last three methods are the same for both proteins. Similarly, in the case of α -synuclein no clear trend can be seen based on either time of introduction or the underlying principles of the methods. Potenci, the newest database derived RCCS predictor finishes first, but the oldest small peptide-base method is the next in line.

The importance of pH effects was highlighted in the comparison of RCCS predictors by the SRD-CRRN method even for folded ubiquitin. As some sort of pH correction is available in four of the eight studied predictors, we intended to investigate the effect of pH *via* SRD calculations. For this purpose, we used chemical shift data for α -synuclein, acquired at various pH and temperature values (BMRB: 18857). We classified the amino acid residues as titratable (including aspartic and glutamic acid) and all others as non-titratable. This way, according to the BMRB dataset out of the possible 140 residues 133 are assigned, yielding 22 titratable and 111 non-titratable residues in the 2.16–7.51 pH range, at 283 K. For the analysis we selected representative pH values of 2.16, 4.21 and 7.51. This enables us to see how the different pH-correction schemes deal with a pH value at which most of the titratable side chains are partially protonated, and what happens if the pH is close to the Asp, Glu sidechain pK_a values. In order to avoid unnecessary outliers in SRD-CRRN calculations, at each pH we use only the suitable RCCS predictors. This means, for example, that the Schwarzsinger method is excluded at pH = 7.51, as its RCCSs have clearly been calibrated under acidic conditions. Similarly, ncIDP and the methods of Wishart and Wang are excluded at pH = 2.16 values, as the Wishart dataset corresponds to pH = 5.0, while data recorded between pH values of 4.0 and 7.5 were used for the development of ncIDP. The Wang method was also developed for very mildly acidic and close to neutral conditions. The obtained pH-dependent SRD plots are shown in Fig. 10.

Considering the titratable residues of α -synuclein at pH = 2.16 the SRD results suggest that the Kjaergaard and Potenci methods – the two RCCS predictors with continuous pH correction schemes – perform the best. Quite surprisingly, the Schwarzsinger method ends up in the in significant range and is the worst at reproducing the consensus of the five methods even under these conditions. Since the pH in this case is rather close to the calibration pH of the Schwarzsinger dataset, this result is surprising. It is also interesting that the two small peptide methods (Kjaergaard and Schwarzsinger) finish first and last, respectively. This confirms that it is generally not recommended to choose an RCCS prediction method solely based on its origin as datasets with very similar backgrounds can produce very different results under conditions at which both should be equally valid. The lack of consensuality between the Schwarzsinger method and its peers might be explained by the small peptides used for calibration. The simple glycine frame

Table 4 The ANOVA and Bonferroni test results for SRD-CRRN data of α -synuclein, BMRB 27348

p (ANOVA)	Method	Mean SRD score	G1	G2	G3	G4	G5	G6	G7
$<10^{-7}$	Potenci	25.66			****				
	Wishart	30.49	****						
	ncIDP	30.83	****						
	Prosecco	31.65		****					
	Kjaergaard	34.19				****			
	Schwarzsinger	40.81					****		
	Wang	41.87						****	
	Camcoil	45.32							****



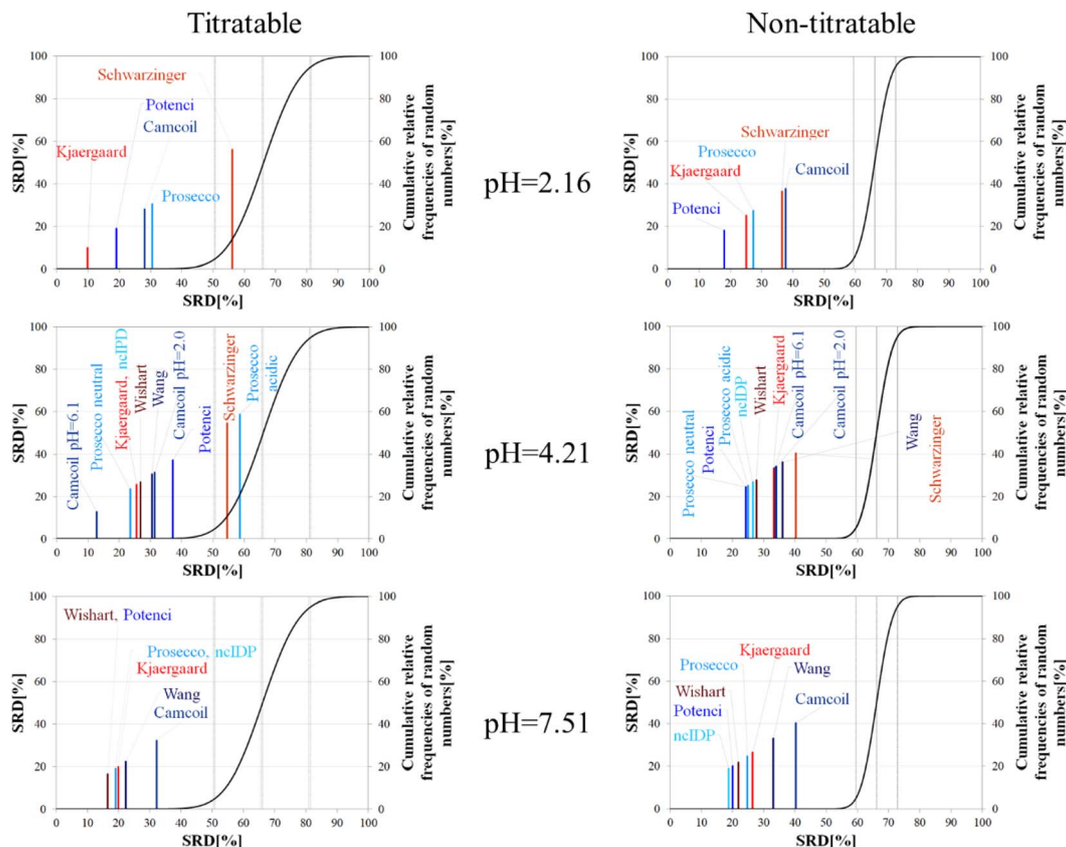


Fig. 10 SRD-CRRN results for C^α SCS vectors for the titratable and non-titratable residues of α -synuclein (BMRB 18857) at different pH values and $T = 283$ K. Small-peptide RCCS predictors (Wishart, Schwarzing, Kjaergaard) are shown in shades of red, while database-derived ones (Wang, Camcoil, ncIDP, Prosecco, Potenci) are shown in shades of blue.

harboring a glutamic or aspartic acid residue is extreme with respect to molecular mobility and intramolecular electronic interactions.

Regarding the non-titratable residues of α -synuclein at pH = 2.16, the order of the same five RCCS predictors is shuffled, and the grouping pattern is also different. In this case, Potenci is the most consensual of the five, while Camcoil finishes last. The Schwarzing method is at the fourth position but is better than random numbers, indicating that the Schwarzing method predicts more consensual RCCSs for non-titratable residues than for titratable ones under acidic conditions. The Kjaergaard method, which was the most consensual for the titratable residues, now finishes second with Prosecco closely following it. Interestingly, at pH = 2.16, all the above differences in SRD values are significant (Tables S1 and S2[†]).

At the intermediate pH of 4.21 and for titratable residues neutral generally neutral methods perform well, while the Schwarzing method and the acidic version of Prosecco end up in the insignificant region (Fig. 10). The most interesting feature at this pH is the very pattern of the SRD-CRRN plots with titratable and non-titratable residues. In the latter case, all methods are grouped close to one another. This highlights that differences between RCCS predictors are generally much more pronounced for glutamic acid and aspartic acid residues than for the non-titratable amino acids. Note, that the general

consensus of the predictors is generally worse at this pH, indicated by higher values SRD values. Also, at pH = 4.21, more predictors perform equivalently well based on *post hoc* Bonferroni test results shown in Tables S3 and S4.[†] Here, there are multiple homogenous groups containing more than one predictor.

For titratable residues at pH = 7.51, the Wishart and Potenci methods perform equivalently and are followed rather closely by the equivalently performing Prosecco, ncIDP and the Kjaergaard method (Table S5[†]). The Wang method is reasonably close to this cluster, while Camcoil is at the seventh position. Once again, Potenci as the newest database-derived method – shows similarity to the Wishart dataset which is the oldest of those used here and has been calibrated on a set of small peptides. It is also interesting that Potenci and ncIDP, where the former is an enhanced and more complete version of the latter, are not the closest to each other. Regarding the non-titratable residues at pH = 7.51 (Table S6[†]), ncIDP seems to be the most consensual, closely followed by Potenci and the Wishart method. Then, Prosecco and the Kjaergaard method perform rather similarly to each other with the Wang method and Camcoil loosely following them.

Generally, the consensus of RCCS predictors is weaker for Glu and Asp residues of α -synuclein, than for non-titratable



ones. The identity of the most consensual predictors is dependent both upon pH and amino acid constitution of the protein.

Besides pH, the other dominant factor affecting CSs and corrected for by some RCCS prediction methods is the temperature. To study its effect, we used the five datasets of BMRB entry 18 857 containing the experimental CSs of α -synuclein at 278, 288, 293, 298 and 303 K, at pH = 5.87. As the Schwarzingger method has been shown to be inappropriate at this pH especially for glutamate and aspartate residues making up more than 10% of amino acids in α -synuclein, we excluded this predictor from the analysis. The results of the SRD calculations for each temperature are shown in Fig. 11. The general pattern shows that ncIDP is the most consensual followed by a group comprising Potenci, Prosecco, and the methods of Wishart, Kjaergaard and Wang in varying order. Camcoil ends up at the seventh position in all cases. Most methods are usually significantly different from each other according to the Bonferroni *post hoc* test, however, there is usually a single pair of methods which are indistinguishable (Tables S7–S11†). At 278 K it is Potenci and Prosecco, at 288 K the Kjaergaard and Wang methods, at 293 and 298 K it is Prosecco and the Wishart method, while finally, at 303 K ncIDP and Potenci end up being indistinguishable at a 5% level of significance. We note that this set of methods shows considerable stability in SRD, namely, at 288 K and above the last three positions are occupied by the

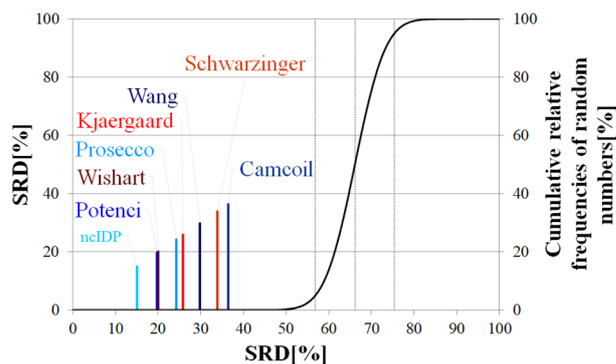


Fig. 12 SRD-CRRN results for C^α SCS vectors of p53TAD¹⁻⁶⁰. Small-peptide RCCS predictors (Wishart, Schwarzingger, Kjaergaard) are shown in shades of red, while database-derived ones (Wang, Camcoil, ncIDP, Prosecco, Potenci) are shown in shades of blue.

method of Kjaergaard, Wang and Camcoil in this exact order. Prosecco and the method of Wishart are close in all cases, switching places between 293 K and 298 K and then their earlier order is restored again at 303 K. Potenci, which performs ordinarily at 278 K, becomes very closely the most consensual method at 303 K.

Since the changes in the experimental CSs of α -synuclein are apparently not related to any change in structural propensities

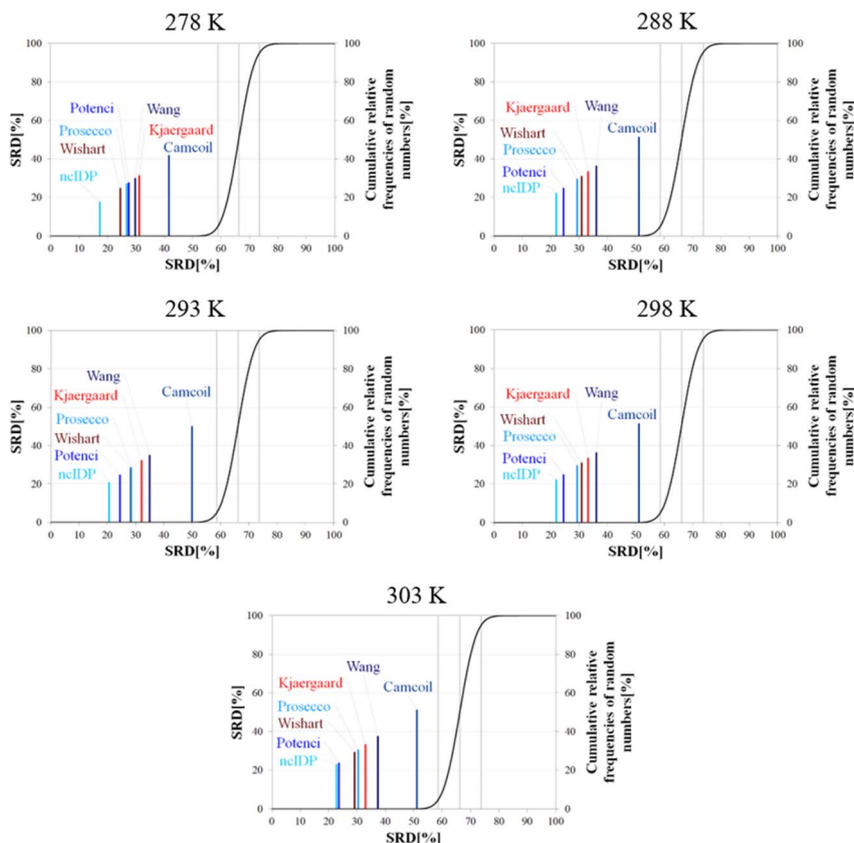


Fig. 11 SRD-CRRN results for C^α SCS vectors of α -synuclein (BMRB 18857) at different temperatures and pH = 5.87. Small-peptide RCCS predictors (Wishart, Schwarzingger, Kjaergaard) are shown in shades of red, while database-derived ones (Wang, Camcoil, ncIDP, Prosecco, Potenci) are shown in shades of blue.



Table 5 The ANOVA and Bonferroni test results for SRD-CRRN data of p53TAD¹⁻⁶⁰

<i>p</i> (ANOVA)	Method	Mean SRD score	G1	G2	G 3	G4	G5	G6	G7
<10 ⁻⁷	ncIDP	15.35		****					
	Potenci	19.92	****						
	Wishart	20.20	****						
	Prosecco	24.49			****				
	Kjaergaard	26.01				****			
	Wang	30.05					****		
	Schwarzinger	34.21						****	
	Camcoil	36.51							****

in this temperature range, the data further highlight the fact that the degree of similarity of RCCS datasets provided by the different predictors is much dependent on the experimental conditions. This is true even despite some general tendencies in the SRD plots displaying stability. The individual changes in the consensuality of the seven methods indicates that in order to formulate a sound conclusion from SCS data, the simultaneous use of a carefully selected set of RCCS predictors is desirable.

3.2.4. Consensuality of RCCS predictors for p53TAD¹⁻⁶⁰: the case of intermediate disorder. As different IDPs behave differently, in order to test our conclusions, we investigated the behavior of another IDP, the p53TAD¹⁻⁶⁰ which has been shown to bear helical propensities in certain regions (Phe19–Lys24, Asp41–Leu43 and Asp49–Glu51). Based on the CS values determined in our earlier work we performed the statistical analysis for C^α SCS data (Fig. 12) The resulting general pattern is similar to that obtained for α -synuclein. However, the SRD values are slightly lower than for α -synuclein, showing a generally stronger consensus among the RCCS predictors, and this is due to the fact, that p53TAD¹⁻⁶⁰ has more pronounced residual structural motifs than α -synuclein. Still, most predictors are shown to be statistically different by the Bonferroni *post hoc* tests (Table 5), Potenci and the Wishart method form the only exception. There is, once again, no clear trend with respect to the theoretical basis or the age of the methods. As in various earlier cases, the Wang, Schwarzinger and Camcoil methods occupy the last three spots, while the Wishart method finishes among the most recent database-derived approaches: ncIDP, Potenci and Prosecco.

4. Conclusions

NMR chemical shifts and the calculated SCSs are the most important atomic-scale variables reporting on the secondary structural propensities of IDPs. This information can be further used for building databases of short linear motifs, and highlighting these regions is important in further interaction studies. On the other hand, SCS analysis is becoming a cornerstone for the possible targeting of IDPs and drug development. However, the SCS method suffers from the ambiguity of RCCSs which makes finding the proper RCCS values an important issue especially in IDP studies. Because of the continuous increase in the number of existing predictors, resulting from both new theoretical approaches and the growth of available

experimental data in the field, RCCS prediction requires precaution when applying the SCS method.

In this work we intended to give an overview with corresponding theoretical background of RCCS prediction, and special attention was paid to neighbor-, pH-, and temperature correction schemes. We attempted summarizing earlier work in a way that highlights fundamentals of different approaches and the temporal evolution of RCCS predictors in the last decades. We believe, such a review might be helpful for the experimental protein scientists to maximally exploit the acquired data for identifying secondary structural propensities of IDPs. Our approach of treating RCCS prediction as a somewhat ill-defined calibration problem is expected to give hints for computational and theoretical researchers for making developments in RCCS prediction.

Our selected set of eight RCCS predictors (most of them are also the ones most abundantly referenced in publications) can, in our opinion, be plausibly used. Performing a visual and statistical comparison of the selected RCCS predictors demonstrates how the choice of any single RCCS predictor might affect the structural conclusions. Even though we focus only on the C^α environment, and the conclusions reflect the behavior of this atom type, the chosen examples highlight general tendencies and are not compromised in validity. We introduce in this field and suggest the use of the very sensitive SRD-CRRN analysis coupled with *post hoc* complemented ANOVA. This approach can detect statistically significant differences even in the case of a folded protein with well-characterized structure. However, these differences only slightly affect secondary structural conclusions, as we show on the example of ubiquitin. Much more importantly, using α -synuclein and p53TAD¹⁻⁶⁰ as examples, we demonstrate that the slight differences – occurring exclusively as a consequence of choosing different RCCS predictors – can be crucial in the study of IDPs. The non-equivalence of RCCS predictors could clearly highlight or mask certain potential secondary structural tendencies. Relying on the pure review part of the present work, we discuss how these differences of RCCS predictors are related to the different theoretical backgrounds of the predictors, especially correction schemes and the molecular systems used in the calibration processes. These examples and explanations were aimed at highlighting the most general pitfalls to avoid during SCS analysis. We have demonstrated that amino acid sequence, pH and temperature mutually determine which RCCS predictors



should be used. However, a trustworthy selection of RCCS predictors should be based on statistical analysis, rather than intuition or habit alone. Especially, applying SRD-CRRN – one of the up-and-coming tools of chemometrics – for SCS analysis and selection of RCCS predictors is an important methodological advance.

In case of IDPs the use of multiple RCCS predictors is beneficial. Noting their individual drawbacks and peculiarities, we recommend all eight predictors used in this study, if experimental conditions permit. However, we believe that there is a need for the development of at least one composite statistical method which is capable of incorporating the information content of multiple RCCS predictors. Until the underlying problem of ill-defined RCCS calibration is satisfactorily solved, such an approach to SCS analysis would be the best and most user-friendly option in the field.

Author contributions

Both D. K. and A. B. contributed to the writing of this manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

Financial support of the National Research, Development and Innovation Office, Hungary (grants NKFI K124900 and K137940) is highly acknowledged.

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