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## What are the minimal folding seeds in proteins? Experimental and theoretical assessment of secondary structure propensities of small peptide fragments<sup>+</sup>

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Certain peptide sequences, some of them as short as amino acid triplets, are significantly overpopulated in specific secondary structure motifs in folded protein structures. For example, 74% of the EAM triplet is found in  $\alpha$ -helices, and only 3% occurs in the extended parts of proteins (typically  $\beta$ -sheets). In contrast, other triplets (such as VIV and IYI) appear almost exclusively in extended parts (79% and 69%, respectively). In order to determine whether such preferences are structurally encoded in a particular peptide fragment or appear only at the level of a complex protein structure, NMR, VCD, and ECD experiments were carried out on selected tripeptides: EAM (denoted as pro-' $\alpha$ -helical' in proteins), KAM( $\alpha$ ), ALA( $\alpha$ ), DIC( $\alpha$ ), EKF( $\alpha$ ), IYI(pro- $\beta$ -sheet or more generally, pro-extended), and VIV( $\beta$ ), and the reference  $\alpha$ -helical CATWEAMEKCK undecapeptide. The experimental data were in very good agreement with extensive quantum mechanical conformational sampling. Altogether, we clearly showed that the pro-helical vs. pro-extended propensities start to emerge already at the level of tripeptides and can be fully developed at longer sequences. We postulate that certain short peptide sequences can be considered minimal "folding seeds". Admittedly, the inherent secondary structure propensity can be overruled by the large intramolecular interaction energies within the folded and compact protein structures. Still, the correlation of experimental and computational data presented herein suggests that the secondary structure propensity should be considered as one of the key factors that may lead to understanding the underlying physico-chemical principles of protein structure and folding from the first principles.

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## 1. Introduction

Understanding fully the relation between the amino acid sequence and the three-dimensional structure of proteins has been a subject of intense research over the last six decades.<sup>1-3</sup> The recent unprecedented success of DeepMind's AlphaFold2 (AF2) algorithm at the 14th Critical Assessment of Structure

Prediction (CASP14)<sup>4</sup> contest is viewed as a major breakthrough in predicting protein 3-D structures.<sup>5</sup> However, deep-learning neural networks used in AF2 do not reveal too many of the underlying physico-chemical/biophysical principles. In fact, the process by which AF2 and other state-of-the-art algorithms reach the final protein structure does not necessarily correspond to the steps of actual protein folding as described by experimental studies.<sup>6</sup> Thus, there is still a need for a deeper understanding of the 'Aufbau' principle of protein 3-D structures and protein folding by an *ab initio* approach. This may allow us to fully grasp the beauty of one of nature's most fundamental processes.

The traditional physical chemist's view of protein folding acknowledges a delicate interplay between several enthalpic and entropic terms, including interactions of the protein surface with the environment (solvent). On the protein side, the enthalpic contributions can be decomposed into an (unfavorable, destabilizing) local strain energy and mostly favorable (stabilizing) intramolecular (inter-residual) interaction energy. Strain energy appears because small fragments of the protein

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<sup>†</sup> Electronic supplementary information (ESI) available: Tables S1–S9 and Fig. S1–S26, including an in-depth discussion of various experimental details, primary computational data (SI\_geoms\_energies.zip file containing all the coordinates of the final QM-optimized peptide structures with their absolute DFT-D3//COSMO-RS energies in methanol), and the XLSX spreadsheet with  $\Delta G_{\rm HPPII}$  values for all 8000 tripeptides extracted from ref. 22. See DOI: https://doi.org/10.1039/d3sc04960d

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are not in their optimal geometry. We have shown that the strain energy may easily reach up to ~5 kcal mol<sup>-1</sup> per amino acid residue<sup>7</sup> and is then expected to be compensated by the favorable intramolecular interactions. Interestingly, it seems that it is rather the favorable intramolecular interaction than low strain, which is conserved by evolution.<sup>7,8</sup> Indeed, it has already been demonstrated that Flory isolated-pair hypothesis is invalid due to the significant interactions between neighboring amino acids.<sup>9-11</sup> Since proteins exist in the condensed phase, the solvation (free) energy difference between the folded and unfolded states of a protein also plays a huge role in determining the final structure.<sup>12</sup> Last but not least, the changes in the solvent entropy as well as the reduction of the conformational entropy of the protein are also considered to be major factors in its folding and stable conformations.<sup>13-16</sup>

One of the key questions – related to the above physicochemical principles – that remains largely unsolved is whether the determinants of a secondary structure are "imprinted" in shorter protein building blocks, *i.e.* polypeptide chains of varying lengths.<sup>17–21</sup> Do the polypeptide chains comprising proteins have variable 'stiffness' that predetermines them to be preferably used in one or the other secondary structure motif? Or is the protein structure a purely global phenomenon that only appears at the level of the full-length sequence of a protein?

To address this question, we recently presented a series of computational and bioinformatics studies providing a more rigorous theoretical framework to address protein folding from first principles (*ab initio*).<sup>7,8,22–24</sup> First, for each of all 8000 possible canonical amino acid triplets ( $X_1X_2X_3$ ), we evaluated statistical probability of finding  $X_1X_2X_3$  in a particular

secondary structure motif (mostly helical or extended) in any protein in a non-redundant subset of the Protein Data Bank (Top8000 database).<sup>23</sup> This allowed us to identify the statistically most pro-helical ( $\alpha$ -helix) and pro-extended (*i.e.*, torsion angles corresponding to a single strand of the  $\beta$ -sheet) amino acid triplets (Fig. 1). Populations on both ends of the helical/ extended 'distribution' were close to 80% which we consider statistically significant (*e.g.*, EAM triplet is found 74% in  $\alpha$ -helical, 3% in extended, and the rest mostly in unstructured parts of proteins, whereas VIV is found 79% in extended and 8% in  $\alpha$ -helical).

We correlated this statistically observed propensity with the results of a large-scale quantum mechanical conformational study on the corresponding N- and C-termini capped tripeptides.<sup>22</sup> The computed free energy differences between the lowest-energy helical and extended conformers of the capped tripeptide, N-Ac-X<sub>1</sub>X<sub>2</sub>X<sub>3</sub>-NHCH<sub>3</sub>,  $\Delta G_{HE} = G(\text{lowest helical}) -$ G(lowest extended), showed that pro-helical tripeptides (such as EAM) tend to have lower  $\Delta G_{\text{HE}}$  values, by 1–2 kcal mol<sup>-1</sup>, than pro-extended ones (such as VIV).23 Thus, they might be considered more suitable building blocks for a-helices than their pro-extended counterparts (and vice versa), which is in line with their populations in protein secondary structures (vide supra). This suggested that the propensities for adopting a particular secondary structure might indeed be encoded in short peptide fragments. In addition, we showed on a limited set that the 'pro-extended' tripeptides/triplets benefit from the presence of an interacting partner to a significantly greater degree than the 'pro-helical' triplets.23

In this work, we materialized our theoretical findings and computational predictions by synthesizing selected (capped)



Fig. 1 Secondary structure preferences of selected pro- $\alpha$ -helical and pro-extended ( $\beta$ -sheet) amino acid triplets in the Top8000 subset of the PDB. The original analysis<sup>23</sup> was updated using the DSSP algorithm, version 4.3,<sup>25,26</sup> which can also detect polyproline II helices. Only triplets where all three amino acids adopt the same secondary structure were considered, *i.e.*,  $\alpha\alpha\alpha$  for  $\alpha$ -helix,  $\beta\beta\beta$  for  $\beta$ -sheet, *etc.* "Mixed" triplets, such as  $\alpha\alpha\beta$  and unordered structures were included in the category "Other". Bend, bridge, and  $\pi$ -helix secondary structures were detected in less than 1% of cases for all selected triplets.

tripeptides with expected extended or helical propensities. We do not expect that the short peptide sequences would adopt a single conformation or would form stable helices (though *N*-Ac-X<sub>1</sub>X<sub>2</sub>X<sub>3</sub>-NH<sub>2</sub> species have exactly a minimal length for one  $\alpha$ -helical turn) or purely extended forms. However, we may expect to find some tendencies (propensities) to one or the other type of secondary structures. For this aim, we probed their structural features experimentally, combining nuclear magnetic resonance (NMR) and circular dichroism (vibrational – VCD and electronic – ECD) spectroscopies. These are excellent, and to a certain degree complementary, tools for gaining valuable insights into the structure of biomolecules in solution.<sup>27-32</sup>

There are several NMR observables affected by the conformation of peptides: chemical shifts, indirect couplings (*J*couplings), temperature dependence of chemical shifts of amide hydrogens or the nuclear Overhauser effect.<sup>33-35</sup> Our investigation of the secondary structure with NMR is mostly based on the measurement of temperature dependence of the  ${}^{3}J_{\rm NH,H\alpha}$  coupling constants. Indirect coupling (*J*-coupling) has become an indispensable NMR parameter for structural analysis because it is closely related to molecular conformation according to the Karplus equations.<sup>36-41</sup> The relation between amide NH and H $\alpha$  hydrogen atoms  ${}^{3}J_{\rm NH,H\alpha}$  (in Hz) and the backbone torsion angle  $\varphi$  has been calibrated on known structures:<sup>42</sup>

$${}^{3}J_{\rm NH,H\alpha} = 6.4\cos^{2}(\varphi - 60^{\circ}) - 1.4\cos(\varphi - 60^{\circ}) + 1.9$$
 (1)

As a rule of thumb, helices exhibit  ${}^{3}J_{\rm NH,H\alpha}$  lower than 6 Hz,  $\beta$ sheet structures exhibit  ${}^{3}J_{\rm NH,H\alpha}$  higher than 8 Hz and random coil structures are in between.<sup>27</sup> An advantage of *J*-couplings is that they are not significantly dependent on solvent<sup>43</sup> or temperature,<sup>44,45</sup> *i.e.* any temperature dependence of *J*-couplings most probably reflects a conformational change. The temperature dependence of  ${}^{3}J_{\rm NH,H\alpha}$  was recently used in a study of short peptides and was interpreted in terms of conformational redistribution.<sup>46</sup>

A disadvantage of NMR is that only the  $\varphi$  angle of the Ramachandran plot could be measured (on non-labeled peptides) and thus the technique may not distinguish left-handed polyproline II (PPII) and right-handed ( $\alpha$ -) helices. Information about the  $\psi$  angle (to distinguish between PPII and  $\alpha$ -helix) can be obtained from NMR experiments with <sup>13</sup>C and <sup>15</sup>N-labeled peptides.<sup>47–49</sup> However, PPII conformation is mostly found in unordered peptides, while it is rarer in proteins (*c.f.* Fig. 1 and also ref. 50). The helical chirality can be well distinguished by CD spectroscopy (VCD or ECD), which, however, does not provide residue-specific information, distinguishing (*e.g.*)  $\alpha\beta\beta$  *vs.*  $\beta\beta\alpha$  conformations. Instead, CD spectra reflect the average conformation.<sup>28,32</sup>

The experimental data for all studied peptides were complemented by accurate quantum chemical calculations including the solvation (DFT-D3//COSMO-RS), calibrated in the previous work.<sup>51</sup> These followed exhaustive conformational sampling covering all three structural motives and provided unambiguous structure/energy mapping. The correlation of experimental and theoretical data allowed us to make several conclusions concerning the bottom-up approach in protein structure predictions *ab initio*.

### 2. Methods

#### 2.1. Selected peptides

Based on our previous work,<sup>22–24</sup> we selected five tripeptides with quite pronounced statistical preference for a particular secondary structure in proteins: EAM( $\alpha$ -helical), KAM( $\alpha$ ), ALA( $\alpha$ ), IYI( $\beta$ -sheet/extended), and VIV( $\beta$ ), gauged by their respective secondary structure populations in the threedimensional protein structures (Fig. 1).

In addition, we analyzed the computational data from our previous work.<sup>22</sup> Within the set of all 8000 tripeptides (200 conformers each, comprising the P-CONF\_1.6M database), we ranked the tripeptides by the lowest computed  $\Delta G_{\text{HE}}$  (primary criterion) and  $\Delta G_{\text{H/PPII}}$  (secondary criterion) values. Thus, we searched for the potentially most pro- $\alpha$ -helical tripeptides (*c.f.* SI.xlsx Table (ESI†) with the  $\Delta G_{\text{HE}}$  and  $\Delta G_{\text{H/PPII}}$  values for all 8000 tripeptides). We excluded the tripeptides containing proline, as they are not expected to adopt extended conformations. Also, we preferred to avoid histidines due to their ambiguous protonation states. This resulted in addition of two tripeptides with potential  $\alpha$ -helical propensity: DIC( $\alpha$ ) and EKF( $\alpha$ ). Thus, judged purely from quantum chemical computations, they should belong to the tripeptides with the highest tendencies/propensities for  $\alpha$ -helical structures.

Throughout computations, all peptides were in their most frequent protonation state at pH 7 in water, *i.e.*, K (Lys) and R (Arg) side chains are positively charged, and E (Glu) and D (Asp) side chains are charged negatively. In addition, EAM and IYI tripeptides were also used for the determination of the effect of solvent on their secondary structure (*c.f.* Fig. S10 in the ESI<sup>†</sup>).

For both computational and experimental analyses, we used a model of a peptide with an acetylated N-terminus and amidated C-terminus, shown in Fig. 2.

Finally, a reference CATWEAMEKCK undecapeptide, in which the EAM triplet is in the core of the  $\alpha$ -helix as found in the chain B of the 20- $\alpha$ -hydroxysteroid dehydrogenase (PDBID 1Q5M, Fig. S1 in the ESI†), was investigated.<sup>52</sup> We presumed that it might also adopt a stable  $\alpha$ -helical conformation in solution. As discussed below, this assumption was later confirmed in this study, by both NMR and VCD.

#### 2.2. Experimental

2.2.1 Peptide synthesis. The studied peptides (*N*-Ac-X<sub>1</sub>X<sub>2</sub>X<sub>3</sub>-NH<sub>2</sub>) were assembled in a solid-phase synthesizer Liberty Blue (CEM, USA) by stepwise coupling of the corresponding Fmocamino acids to the growing chain on Rink Amide MBHA resin (100–200 mesh, 0.67 mmol  $g^{-1}$ ) purchased from IRIS, Biotech GmbH, Marktredwitz, Germany. Fully protected peptide resins were synthesized according to a standard procedure involving cleavage of the N $\alpha$ -Fmoc protecting group with 20% piperidine in DMF and coupling, mediated by mixtures of coupling reagents DIC/Oxyma in DMF. On completion of synthesis, the deprotection and detachment of linear peptides from the resins



Fig. 2 N-Acetylated tripeptides used for the calculations and experiments, with the main chain dihedral angles ( $\varphi$  and  $\psi$ ) highlighted.

were carried out simultaneously using a TFA/H<sub>2</sub>O/TIS (95:2.5:2.5) cleaving mixture. Each of the resins was washed with DCM, and the combined TFA filtrates were evaporated at room temperature. The precipitated residues were triturated with *tert*-butyl-methylether, collected by suction, and dried by lyophilization. The linear peptides were purified by HPLC using a Waters instrument with a Delta 600 pump, and a 2489 UV/VIS detector. The purity and identity of all peptides were determined by analytical HPLC and by the ESI MS technique.

2.2.2 NMR experiments. Variable-temperature NMR spectra were recorded on a 500 MHz NMR spectrometer Bruker Avance II<sup>TM</sup> HD (<sup>1</sup>H at 500 MHz, <sup>13</sup>C at 126 MHz) in DMF-d<sub>7</sub> and CD<sub>3</sub>OH for solutions of approximately 1 mg of the peptide in  $600\ \mu L$  of the solvent. Proton spectra were referenced to the solvent signals  $\delta = 2.75$  and  $\delta = 3.31$ , respectively. Proton spectra of CD<sub>3</sub>OH solutions were recorded with presaturation of the intense OH signal. The characterization spectra of the prepared oligopeptides were recorded on a 500 MHz NMR spectrometer Bruker Avance III<sup>TM</sup> HD (<sup>1</sup>H at 500 MHz, <sup>13</sup>C at 126 MHz) or on a 600 MHz NMR spectrometer Bruker Avance III<sup>TM</sup> HD (<sup>1</sup>H at 600 MHz, <sup>13</sup>C at 151 MHz) in DMSO- $d_6$  ( $\delta = 2.50$  (<sup>1</sup>H) and  $\delta = 39.70 \, (^{13}C)$ , DMF- $d_7 \, (\delta = 2.75 \, (^{1}H)$  and  $\delta = 29.76 \, (^{13}C)$ ) or methanol- $d_4$  ( $\delta = 3.31$  (<sup>1</sup>H) and  $\delta = 49.00$  (<sup>13</sup>C)). Complete signal assignment is based on homo- and heteronuclear correlation experiments COSY, TOCSY, ROESY, HSQC and HMBC. The solvents used were purchased from Eurisotop.

2.2.3 VCD experiments. Prior to VCD experiments, TFA remaining from the peptide synthesis was removed according to the published procedure.<sup>53</sup> The purified peptides were dissolved in MeOH (HPLC grade, VWR) at concentrations varying between 2 and 9 mg mL<sup>-1</sup> (5 mM to 20 mM), depending on the solubility. The solutions were placed in a sealed BaF<sub>2</sub> cell with a pathlength of 200  $\mu$ m (International Crystal Laboratories, Inc., Garfield, USA). The VCD and IR spectra were recorded with a ChiralIR-2X VCD spectrometer (BioTools, Inc., Jupiter, USA) for 15 hours with a resolution of 8 cm<sup>-1</sup> at room temperature. Spectra of the solvent (MeOH) recorded under identical conditions were subtracted from the sample spectra and the resulting spectra were subjected to a baseline correction.

**2.2.4 ECD experiments.** The electronic circular dichroism (ECD) measurements were performed with a Jasco-1500 spectropolarimeter equipped with a Peltier thermostatted holder PTC-517 (JASCO, Easton, MD, USA). Tripeptides were dissolved in MilliQ water or in methanol (MeOH) at concentration 1 mg

 $\rm mL^{-1}$ . ECD spectra were measured at room temperature using the following experimental setup: spectral range 195–280 nm, a rectangular quartz cell with path length 0.5 mm, standard instrument sensitivity, 1 nm bandwidth, a scanning speed of 10 nm min<sup>-1</sup>, a response time of 8 s, and one accumulation. The temperature dependencies were recorded only for aqueous solutions in a temperature range from 5 °C to 90 °C with the same experimental setup. The solvents used were purchased from Sigma-Aldrich. Numerical analysis of the secondary structure and secondary structure assignment was performed using the CONTIN program within the CDPro software package.<sup>54</sup>

#### 2.3. Theoretical

2.3.1 Peptide conformer sets. Capitalizing on our previous experience in generating extensive sets of peptide conformers,<sup>22,24</sup> we used the CREST program<sup>55</sup> (Conformer Rotamer Ensemble Sampling Tool, ver. 2.12). CREST runs an iterative search for conformers, involving multiple molecular dynamics, metadynamics, semiempirical optimizations, and semiempirical single point calculations. As commonly used force fields quite often overestimate the population of  $\alpha$ helix,49,56-59 we used the GFN-2 semiempirical QM method60 for optimization and the ALPB implicit solvation model61 with methanol as solvent. Since we consider the accuracy of the GFN-2 single point energies insufficient, we re-calculated the DFT single point energy of the GFN-2 optimized conformers (using the "-xnam" flag of the CREST command line input for such purposes). This calls the external DFT single point calculation, in our case performed using TURBOMOLE, version 7.6.62 We employed the BP86 functional,63 DGauss-DZVP basis set64 and Grimme's D3(BJ) dispersion correction with special parameters for proteins.<sup>65,66</sup> Solvation effects (within the DFT framework) were computed by employing the COSMO (conductor-like screening model)67 and COSMO-RS (COSMO for realistic solvation)68 solvation models as implemented in the BIOVIA COS-MOtherm 2021 program. The "BP\_TZVPD\_FINE\_21.ctd" parametrization file with FINE cavities<sup>69</sup> was used. Final free energies of conformers were obtained via the following formula:

$$G = E_{\rm COSMO} + \Delta E + \mu \tag{2}$$

where  $E_{\text{COSMO}}$  corresponds to BP86-D3BJ/COSMO( $\varepsilon = \infty$ ) energy of the molecule,  $\Delta E$  is the averaged correction for the dielectric

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energy, and  $\mu$  is the chemical potential of the conformer. As inherent in the COSMO-RS procedure, 'scaling' from an ideal conductor to the real solvent with a given permittivity is included in the  $\Delta E$  and  $\mu$  terms. All these values were provided by the COSMOtherm program (version 21). As a last step, we removed redundant conformers using the same approach as in our previous work,<sup>22</sup> but this time applied only to backbone dihedral angles, ignoring the side chain conformations.

**2.3.2 Explicit solvation.** It has been shown that both conformational changes and different secondary structure equilibria in more hydrophobic peptides are strongly affected by hydrogen bonds between the solute and solvent molecules.<sup>16</sup> The same holds for the frequencies and intensities in IR and VCD spectra.<sup>70,71</sup> Both of these illustrate the importance of explicit solvation as already published.<sup>72</sup> Therefore, we added a limited explicit first solvation layer using a repeated neighbor search as implemented in the Biopython library<sup>73</sup> for every possible combination of one, two, three, and four N-H···O(H)Me hydrogen bonds that can be formed with backbone amides opposite to the carbonyl oxygen (Fig. 3). The procedure resulted in (maximally, depending on whether there is enough space for solvent molecules) four single-solvated, six double-solvated, four triple-solvated, and one quadruple-solvated structures.

**2.3.3** Calculation of VCD spectra and final energies. For the calculations of VCD spectra, only conformers with relative energies up to 6 kcal mol<sup>-1</sup> from the global minima obtained from extensive conformational sampling (without explicit solvent) were considered. Each conformer was then solvated according to the procedure described above and geometry of clusters was re-optimized using the Gaussian16 program,<sup>74</sup>

employing B3-LYP functional,75,76 6-31+G(2d,p) basis set, D3(BJ) empirical dispersion correction,65,77 conductor-like polarizable continuum model (CPCM),78 and dielectric constant corresponding to methanol ( $\varepsilon_r = 33$ ). This combination has been shown to give good results for very similar peptide fragments.79 Vibrational frequencies and IR and VCD intensities were then calculated at the harmonic level. To estimate Boltzmann population, the methanol molecules were removed from the clusters and single point (free) energies were calculated, according to eqn (2), at the BP86-D3(BJ)//(COSMO-RS) level, employing the def2-TZVPD basis set.<sup>80</sup> The line intensities were extracted from the Gaussian 16 output and convoluted with Lorentzian curves with a bandwidth of 10 cm<sup>-1</sup>. Contribution of methanol to the computed spectrum was removed by deleting the polar and axial tensors of methanol atoms, using our in-house program eattt, as described in ref. 81. Final spectra of all tripeptides were obtained by Boltzmann weighting of conformers, using single point energies. This computational protocol was validated on model alanine tripeptides of pure *α*-helical, extended and PPII conformations (for details see ESI, Fig. S2<sup>†</sup>).

## 3. Results

# 3.1. VCD and ECD spectra of the studied tripeptides in solution and comparison with those of the reference undecapeptide CATWEAMEKCK

For ALA, DIC, EAM, EKF, KAM, and VIV, the VCD and IR spectra are depicted in Fig. 4, whereas ECD spectra can be found in Fig. 5. Due to its poor solubility, we were not able to measure any CD spectrum of the IYI tripeptide.



Fig. 3 An example of quadruple explicit solvation of tripeptide ALA with four methanol molecules.



Fig. 4 Experimental VCD (left) and IR (right) spectra of six capped tripeptides and one undecapeptide in the amide I region measured in methanol. Intensity of the spectra of the CATWEAMEKCK undecapeptide was scaled by 0.27 (approx. 3/11) for easy comparison.

VCD spectra of EAM and VIV show a predominantly negative band in the amide I region at around 1650  $\text{cm}^{-1}$ , with weak positive lobes at  ${\sim}1670~{\rm cm}^{-1},$  and  ${\sim}1630~{\rm cm}^{-1}.$  It is significantly shifted to lower wavenumbers with respect to the IR absorption, which has a maximum at 1670-1675 cm<sup>-1</sup> (see Fig. S3<sup>†</sup> for detailed comparison). Such a pattern implies significant content of  $\beta$ -sheets,<sup>82-84</sup> which could include both extended β-strands and possibly a certain contribution of intermolecular  $\beta$ -sheets occurring due to potential peptide aggregation at high sample concentrations used in the VCD experiments. In particular, the IR band at  $\sim 1622 \text{ cm}^{-1}$  of EAM, typical for intermolecular β-sheets,<sup>85,86</sup> could be connected to the presence of aggregated species in EAM (Fig. S3 in the ESI<sup>†</sup>). The absence of such a band for VIV implies that its VCD spectrum likely comes from its inherent propensity for extended βstrand conformation.

This is consistent with the ECD data obtained at lower sample concentrations minimizing the chance of aggregation. A distinct negative band at around  $\sim$ 220 nm in ECD spectra of VIV (particularly in water) also suggested the presence of a  $\beta$ -sheet in addition to random coil/PPII indicated by the intense negative band at around 197 nm. Therefore, we can assume that the

major conformation of VIV is indeed the extended β-strand. This is consistent with the published values for the similar VVV tripeptide: 68% of the  $\beta$ -strand secondary structure with the remaining contributions from PPII and the α-helix.87,88 In contrast, for EAM we may assume that  $\beta$ -type contribution in its VCD spectrum could come from the intermolecular β-sheet of the aggregated species, or from a combination of an intermolecular  $\beta$ -sheet in aggregated molecules and extended  $\beta$ -strand in non-aggregated ones. A more pronounced negative band at  $\sim$ 1645 cm<sup>-1</sup> and blue-shifted to  $\sim$ 1677 cm<sup>-1</sup> positive lobe common for PPII conformation suggest larger content of PPII structure in EAM, while a weaker negative shoulder at  ${\sim}1658~\text{cm}^{-1}$  might come from a smaller contribution of the  $\alpha\text{-}$ helix.82,84 This assumption is generally corroborated by the ECD data for EAM in methanol, showing largely random coil/PPII conformation with some minor  $\alpha$ -helical contribution (Fig. 5). Thus, PPII,  $\alpha$ -helical and, possibly, extended  $\beta$ -strand secondary structures could be potentially accessible for the EAM tripeptide.

For DIC, which is a tripeptide with one of the lowest  $\Delta G_{\text{HE}}$  values (*ca* -2 kcal mol<sup>-1</sup>, *c.f.* SI.xlsx Table (ESI<sup>†</sup>) and ref. 23), VCD spectra are characterized by a large negative spectral



Fig. 5 ECD spectra of six tripeptides and one undecapeptide measured in methanol and water.

band at ~1660 cm<sup>-1</sup> accompanied by a weak positive shoulder at ~1711 cm<sup>-1</sup> suggesting that it is a combination of  $\alpha$ -helix and PPII, with significantly higher  $\alpha$ -helix content compared

to all other studied tripeptides. While the typical VCD spectrum of the  $\alpha$ -helix is characterized by a positive (-/+) couplet (*c.f.* CATWEAMEKCK peptide in Fig. 4 featuring the

distinctive 1668(-)/1644(+) couplet), we explain in detail the untypical shape of the DIC spectrum in the ESI (Fig. S4)† and discuss it also in Section 3.4 below (comparison of the calculated and experimental VCD). The remaining three tripeptides – KAM, ALA, and EKF – show the highest content of PPII (more visible in cases of ALA and EKF)<sup>82,89</sup> in the VCD spectra characterized by a negative (~1685 cm<sup>-1</sup> (+)/~1655 cm<sup>-1</sup> (-)) couplet typical for this structure. This compares well with the published values<sup>87</sup> suggesting 84% of the PPII secondary structure for AAA and other XXA tripeptides.

The ECD spectra of DIC, KAM and ALA are generally consistent with the VCD data. Similarly to VCD, ECD suggests the highest α-helical propensity for DIC (even in water) and mainly the PPII structure for KAM and ALA in methanol and water. Interestingly, the ECD spectra of EKF show high PPII content in combination with an extended structure and no contribution from the  $\alpha$ -helix in water and methanol (see description in the ESI and Table S1<sup>†</sup> for details). It is worth mentioning that we did not experimentally observe the S-S bond formation between DIC tripeptides. In addition, we also measured the VCD and ECD spectra of the reference CAT-WEAMEKCK undecapeptide. CATWEAMEKCK is the longest αhelix which contains an EAM tripeptide in the middle, found in the Top8000 data set. The undecapeptide exhibits a clear character of α-helix in its VCD spectrum (negative/positive doublet at 1668 cm<sup>-1</sup>(-)/1644 cm<sup>-1</sup>(+)).<sup>82,84</sup> ECD also indicates α-helix, with negative minima at 207 nm and 223 nm.32,54 Therefore, CATWEAMEKCK is an example of an α-helix stable in solution.

# 3.2. NMR spectra of the studied tripeptides in solution and comparison with that of the reference undecapeptide CATWEAMEKCK

To obtain independent, somewhat complementary experimental information, we employed NMR spectroscopy to characterize the structure of the pro-helical ALA, DIC, EAM, EKF, and KAM, and pro-extended VIV and IYI tripeptides in solution (using DMF and methanol as solvents).

Fig. 6 depicts the NH region of variable-temperature <sup>1</sup>H NMR spectra of EAM in methanol whereas the spectra in DMF are shown in the ESI (Fig. S6).<sup>†</sup> For EAM in methanol at room temperature, the <sup>3</sup> $J_{\rm NH,H\alpha}$  coupling values of all three amino acids fall in the range typical for random-coil structures (Table 1) composed by a mixture of helical and extended conformers. However, variable-temperature experiments reveal that the couplings of all three amino acids decrease with decreasing temperature (Table 1), which indicates that the population of helical ( $\alpha$ - or PPII) structures increases at lower temperatures. Similar conclusions can be made from the NMR data obtained in DMF which are deposited in the ESI (Table S3).<sup>†</sup>

The NMR measurements in less polar DMF (Tables S2–S8†) have a slightly different temperature window (360–240 K) but also cover more than a 100 K range. The value of  ${}^{3}J_{\rm NH,H\alpha}$  coupling in the glutamic acid (residue E) in EAM is, at 300 K, similar in both solvents, and the  $\Delta J$  value (the change of the coupling values induced by a 100 K decrease in temperature) is also similar. On the other hand, the  ${}^{3}J_{\rm NH,H\alpha}$  coupling in alanine (residue A) is higher in DMF (6.6 Hz vs. 6.2 Hz in methanol) and the  $\Delta J$  value is significantly lower (–0.6 Hz in DMF vs. –1.0 Hz in methanol). This observation indicates that the propensity of



Fig. 6 The NH region of <sup>1</sup>H NMR spectra of the tripeptide EAM in methanol at T = 200-300 K. The temperature-induced changes in chemical shifts of the signals are caused by an intermolecular exchange of the NH and solvent protons.

Table 1 Experimentally determined  ${}^{3}J_{NH,H\alpha}$  coupling values (Hz) in the ALA, DIC, EAM, EKF, KAM, VIV, and IYI peptides in methanol at T =200-300 K and the change in the coupling values induced by a 100 K decrease in temperature ( $\Delta J_{200-300} = J_{200K} - J_{300K}$ ). For comparison, see the DFT-calculated values of the coupling for ideal  $\alpha$ -helix, extended and PPII conformations in the ESI

T/K	300	280	260	240	220	200	$\Delta J_{200-300}$
ALA							
A1	а	5.8	5.5	5.3	5.1	5.0	$\leq -0.8$
L	а	7.4	7.4	7.2	7.2	7.1	$\leq -0.3$
A3	а	7.0	6.8	6.7	6.3	6.1	≤-0.9
DIC							
D	8.0	8.1	8.1	8.1	8.0	8.0	0.0
Ι	7.1	7.0	6.7	6.6	6.7	6.3	-0.8
С	7.4	7.3	7.1	7.0	6.9	6.7	-0.7
EAM							
Е	6.6	6.3	6.3	6.1	6.0	5.7	-0.9
А	6.2	6.1	6.0	5.7	5.5	5.2	-1.0
М	7.9	7.9	7.8	7.7	7.6	7.5	-0.4
EKF							
Е	6.4	6.3	6.1	5.8	5.6	а	$\leq -0.8$
Κ	7.5	7.4	7.4	7.2	7.1	а	$\leq -0.4$
F	8.0	7.9	7.9	7.8	7.8	7.5	$\leq -0.5$
KAM							
К	7.0	6.9	6.7	6.5	а	6.2	-0.8
Α	6.2	6.1	5.9	5.7	5.5	5.2	-1.0
М	7.8	7.7	7.7	7.6	а	7.2	-0.6
VIV							
$V1^{b}$	7.9	а	а	7.7	7.4	6.9	-1.0
Ι	8.6	а	а	8.6	8.5	а	${\sim}{-0.1}$
$V3^b$	8.7	8.5	8.3	8.3	8.1	8.0	-0.7
IYI							
$\mathrm{I1}^{b}$	8.0	8.1	а	7.7	а		
Y	7.7	8.1	а	7.7	7.8		${\sim}0$
$I3^{b}$	8.7	а	а	8.7	а		

<sup>a</sup> Not determined because of a signal overlap, signal broadening or fast chemical exchange process. <sup>b</sup> The assignment of V1 and V3 in VIV and I1 and I3 in IYI may be interchanged.

the EAM peptide to form some helical structures is higher in methanol than in DMF. The value of the  ${}^{3}J_{NH,H\alpha}$  coupling in methionine is similar in both solvents, and the  $\Delta J$  value is close to zero in DMF, whereas it is -0.4 in methanol. VCD and ECD spectra suggest that the helical conformations observed in EAM by NMR at room temperature are rather of PPII character. Together with the fraction of extended conformations (in VCD mixed with the signal of aggregation), the EAM tripeptide is mostly a combination of all three secondary structure types.

Contrary to the EAM tripeptide, the magnitudes of all  ${}^{3}J_{\rm NH,H\alpha}$ couplings are significantly higher in the pro-extended IYI tripeptide (not measured by VCD) in both solvents (8-9 Hz, Table 1). Furthermore, the  ${}^{3}J_{NH,H\alpha}$  coupling values are almost temperature independent. In DMF, the  $\Delta J$  values can be found between -0.2 and +0.2 Hz. Some of the coupling values in methanol at temperatures below 240 K and at 260 K could not

be obtained because of a signal overlap. However, the coupling values that could be resolved are also almost temperature independent; only the coupling value of one of the isoleucine residues decreased slightly (-0.4 Hz). These characteristics are associated with extended structure motifs; therefore the IYI tripeptide is mostly extended.

Next, we measured the temperature dependence of  ${}^{3}J_{\rm NH,H\alpha}$ couplings in other peptides (ALA, KAM, and VIV) that were previously identified by bioinformatics to have a propensity for the α-helical (ALA and KAM) and extended (VIV) structures. Unfortunately, VIV is poorly soluble in DMF and methanol, and we were not able to obtain the full data set at all investigated temperatures. However, the data that could be obtained clearly show that the  ${}^{3}J_{NH,H\alpha}$  coupling in the central isoleucine residue of VIV is high and almost temperature independent in methanol (Table 1), suggesting mainly an extended structure. The coupling in the valine residues V1 and V3 decreases with decreasing temperature in methanol, which is in line with conformational analysis (vide infra). These results are similar to the published results of the VVV tripeptide in water.<sup>49</sup> Similarly, the  ${}^{3}J_{\rm NH,H\alpha}$  coupling of the central leucine residue in the ALA tripeptide is almost temperature independent. This is different from the statistics in proteins, where L in ALA is mostly in the αhelical conformation. However, the NMR data are in line with the conformational analysis (vide infra). The  ${}^{3}J_{\rm NH,H\alpha}$  couplings and their temperature dependence in the KAM tripeptide are similar to those in EAM. According to the VCD and ECD spectra, helical conformers of KAM are largely of the PPII type (lefthanded helix) and not  $\alpha$ -helical at room temperature.

We also measured the other two tripeptides with computationally predicted propensity towards *α*-helical conformation: DIC and EKF. For DIC, the  ${}^{3}J_{\rm NH,H\alpha}$  coupling in the asparagine residue (D) in methanol is almost temperature independent, while the couplings of the other two amino acid residues are significantly dependent on temperature. Values of these couplings at lower temperature (about 6.5 Hz) point to some form of helical structure (a- or PPII or combination). Similarly, the

Table 2 Experimentally determined  ${}^{3}J_{NH,H\alpha}$  coupling values (Hz) in the residues of CATWEAMEKCK in methanol at T = 320-260 K,  $\Delta\delta NH/\Delta T$ (ppb  $K^{-1}$ ) and chemical shifts of hydrogen atoms H $\alpha$  (ppm, referenced to CD<sub>3</sub>OH,  $\delta = 3.31$ ). Corresponding values for the tripeptide EAM are shown in parenthesis

<i>T</i> /K	320	300	280	260	$\Delta \delta \mathrm{NH} / \Delta T$	$\delta(H\alpha)$
C1	52	5.0	49	47	-65	4 30
A2	4.5	4.6	4.5	4.2	-5.6	4.27
Т	а	а	а	а	а	4.00
W	4.4	4.6	4.5	а	-5.6	4.39
E5	а	4.4 (6.6)	3.8 (6.3)	3.4 (6.3)	-6.4(-5.6)	3.93 (4.28)
A6	4.5	4.6 (6.2)	4.4 (6.1)	4.3 (6.0)	-3.5(-6.7)	4.03 (4.28)
Μ	4.8	4.8 (7.9)	4.7 (7.9)	4.4 (7.8)	-3.7(-6.2)	4.13 (4.43)
E8	а	4.7	4.7	4.4	-3.8	3.97
K9	5.4	5.1	4.9	а	-4.2	4.08
C10	6.7	6.5	6.3	5.9	-1.1	4.30
K11	а	а	а	а	а	4.24

<sup>a</sup> Not determined because of a signal overlap, signal broadening or fast chemical exchange process.

glutamine residue (E) of the EKF tripeptide shows stronger temperature dependence, as the  ${}^{3}J_{\rm NH,H\alpha}$  lowers by 1.0 Hz. The remaining two residues change much less with temperature. The

DIC and EKF tripeptides were also measured in water (H<sub>2</sub>O–D<sub>2</sub>O mixture) at 280 and 300 K (Tables S4 and S5†) and the  ${}^{3}J_{\rm NH,H\alpha}$  coupling constants are similar to those obtained in methanol.



Fig. 7 Histograms of conformer energies for  $\alpha$ -helical, extended, and PPII conformers of seven tripeptides in methanol. Conformer energies were calculated at the BP86-D3(BJ)/def2-TZVPD//COSMO-RS level.

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Lastly, we measured the NMR spectra for the reference CATWEAMEKCK undecapeptide and concluded that it indeed adopts an  $\alpha$ -helix in its EAM core (Table 2, see also Chapter 7 in the ESI† for details), in perfect agreement with the VCD and ECD results presented above.

In addition, we calculated the *J*-coupling values for ideal  $\alpha$ helical, extended, and PPII conformations of all seven tripeptides (see Table S9 in the ESI†), to show that the experimentally determined values fit in the range of the calculated results.

#### 3.3. QM(DFT-D3)//COSMO-RS conformational sampling

In our previous work, a limited sampling of all 8000 tripeptides was performed,<sup>22</sup> employing the calibrated QM protocol.<sup>22,51,66-</sup> However, only 200 initial conformers were generated for each tripeptide, which covered a rather limited part of their vast conformational space. Therefore, we carried out extensive DFT-D3//COSMO-RS//GFN-2 conformational sampling, as described in Methods, of seven selected tripeptides: presumably pro-helical ALA, DIC, EKF, EAM, and KAM, and pro-extended VIV and IYI. This resulted in 608–5179 final conformers for each tripeptide. The results are summarized in Fig. 7, which compares the energetic distribution of  $\alpha$ -helical, extended, and PPII conformers, separately for each amino acid. In this respect, QM calculations can be directly compared to the NMR data discussed above, reflecting the secondary structure of each residue.

The histograms in Fig. 7 illustrate markedly different trends observed among the seven tripeptides. EAM has all three structural types (a-helix, extended, and PPII helix) energetically accessible, which is consistent with the spectroscopic results. DIC exhibits a stronger tendency to form  $\alpha$ -helical structures (with respect to the other peptides studied herein). Moreover, by correlating NMR and computational data on a per-residue basis, we may observe almost perfect agreement between the two. From NMR, the tendency for helicity increases in the order  $D < C \le I$ , which is exactly the case in the DFT-D3//COSMO-RS histograms. The experiments indicated that VIV and IYI prefer extended conformations, and indeed, the VIV and IYI extended conformers are computed to be lower in energy. Furthermore, NMR predicts the tendency for the extended structure in the order  $V_{1/3} < I$  (c.f. Table 1), which is also seen from the computed histograms (Fig. 7). The same holds true for IYI.

Experimentally, EKF and KAM secondary structures seem to be mixtures of PPII helix with minor  $\alpha$ -helix contribution, which is well reproduced by the calculations, both 'globally' and on a per-residue basis. For example, in KAM, the terminal methionine residue has quite a high propensity for extended conformations, which is observed both computationally as well as in NMR. In the case of ALA, NMR predicts that L is assumed to adopt preferably extended conformation, and this can also be seen in computed histograms. Terminal alanine residues behave somewhat differently with respect to each other in NMR (Table 1), which is also observed computationally, as A1 tends to adopt extended conformations less than the A3 residue. We also observed that conformational energy distribution is similar in other solvents, as illustrated in the ESI (Fig. S10)<sup>†</sup> for EAM and IYI. In summary, we demonstrated that predictions provided by quantum chemical calculations are in agreement with the experimentally obtained  ${}^{3}J_{\rm NH,H\alpha}$  coupling constants, VCD and ECD spectral patterns. VCD and ECD spectroscopy nicely complements the NMR experimental data by distinguishing the left- (PPII) and right-( $\alpha$ ) handed helix.

#### 3.4. Theoretical calculations of VCD spectra

We calculated IR and VCD spectra of six tripeptides (ALA, DIC, EAM, EKF, KAM, and VIV; in MeOH). Fig. 8 depicts the amide I region of the six tripeptides. Note that the calculated frequencies in Fig. 8 were shifted down by about 50 cm<sup>-1</sup> to match the



**Fig. 8** Calculated (dashed) and experimental (solid) VCD spectra of capped tripeptides in the amide I region, with methanol as a solvent. Calculated spectra were obtained *via* Boltzmann weighting of spectra of the individual conformers, calculated at the B3-LYP(D3-BJ)/6-31+G(2d,p)/CPCM(methanol) level, using BP86/def2-TZVPD/COSMO-RS energies as weights. Calculated spectra were scaled to fit the experiment, for easy comparison.

experiment. This is a typical computational error arising mostly from limited accounting for the solvent and anharmonic contributions.70,90 It must also be considered that the experimental spectra represent a convolution of spectral patterns characteristic of different structures, as demonstrated in Section 3 of the ESI,<sup>†</sup> and thus cannot be directly assigned to classical spectral characteristic of a single structure. DIC shows the strongest  $\alpha$ -helical character, evident from the negative band at 1660 cm<sup>-1</sup> (scaled spectrum) and only a minor positive signal at higher wavenumbers. Both spectral features agree well with the experiment, where they appear at 1660 and 1711  $\text{cm}^{-1}$ , respectively. EAM exhibits a combination of PPII and α-helix (negative bands at 1645 cm<sup>-1</sup> and at 1679 cm<sup>-1</sup>, with the strong positive band at 1661 cm<sup>-1</sup> coming from the spectral overlap of both these structures; all in the scaled spectrum), pointing out their energetic accessibility. The experimental VCD spectrum of EAM is dominated by the intermolecular  $\beta$ -sheet contribution (coming from partially aggregated peptides in the experiment), with some contribution from PPII, α-helix and possibly an extended β-strand. The ALA tripeptide shows a nearly conservative negative couplet with the negative lobe calculated at  $1655 \text{ cm}^{-1}$  and the positive one at  $1678 \text{ cm}^{-1}$ , which is a typical signature of the PPII structure. The computed spectrum agrees well with the experiment, suggesting a major PPII contribution to this peptide. The calculated spectrum of VIV could be associated with a contribution of PPII and possibly an extended structure (shown by the overall negative signal with a minimum calculated at 1650 cm<sup>-1</sup> and a weaker positive lobe at a higher wavenumber).<sup>82,84</sup> This is in general agreement with the experimental VCD spectrum, illustrating that in VCD, the PPII helix is generally more 'visible' than extended structures, which provide weaker signals.84 Peptides EKF and KAM are mostly a mix of PPII with other secondary structure types, without significant αhelical contribution. This also agrees quite well with the experimental spectra.

## 4. Discussion

Experimental NMR, VCD, and ECD spectra, supported by largescale calibrated66 DFT-D3//COSMO-RS calculations showed that there might indeed be some preference for a particular secondary structure encoded in the peptide fragments as small as tripeptides. These propensities are quite hard to decipher on a complex background given the high conformational flexibility of these small peptide fragments in solution. However, we tried to show that a careful correlation of the experimental (NMR, VCD, and ECD) and computational (DFT) data may represent a strategy to extract the secondary structure propensities. NMR and computations provide detailed local information, which can be decomposed on a per-residue basis, while VCD and ECD spectra are of a more global character. At the current technological level they do not distinguish subtle structural features of individual amino acids within the peptide chain without isotope labelling. In contrast, NMR spectra of (not isotopically labeled) peptides do not distinguish between a- and PPII helices, which is where VCD and ECD spectra provide an important insight. In addition to VCD, variable-temperature

ECD experiments can also distinguish between PPII and random coil conformations (for details see the ESI<sup>†</sup>). We note that PPII is assumed to be more common helical arrangement in shorter peptides, mainly those containing alanine.<sup>49,50,91,92</sup> Also, it has already been shown that for the trialanine residue this does not depend on pH.<sup>93</sup>

Among the studied tripeptides, some were shown to prefer  $\alpha$ -helical arrangement (*e.g.*, DIC), while others, such as VIV and IYI, have inherent propensities for extended conformations. For EAM, the NMR data indicate that there are both extended and helical conformers present, in agreement with the CD spectra which further indicate a PPII helix rather than an  $\alpha$ -helix. Large-scale DFT-D3//COSMO-RS conformational sampling of EAM shows almost equivalent populations of all three secondary structures (incl. PPII). There are also tripeptides with an inherent propensity for PPII, such as ALA, KAM, or EKF; however, they do not preserve this secondary structure in proteins (see Table 1). In fact, PPII conformations are quite rare for the selected triplets in proteins (see Fig. 1).

All of this illustrates that conformational behavior of protein constituents loosely correlates with their (over)populations in a particular secondary structure. This can be traced to fragments as short as tripeptides. For example, EAM and VIV (IYI) tripeptides show a sharp difference in secondary structure preference in proteins ( $\alpha$ -helix/ $\beta$ -sheet, respectively). Our data consistently reproduce the preference of VIV (and IYI) for  $\beta$ sheet conformation on the tripeptide level. Although EAM does not show a clear preference for  $\alpha$ -helical conformers on the tripeptide level, it certainly has a larger tendency toward  $\alpha$ helical conformations than VIV. Thus, some amino acid triplets may "imprint" their accessible (preferred) conformations into the final protein folds. These are by no means "stable" secondary structures, as only some tripeptides exhibit these preferences, while the majority is rather flexible and could be viewed as a model for intrinsically disordered proteins.94 Very importantly, the calculations have shown that the equilibrium between the three (or more) conformational states of tripeptides is very subtle. Energetically, the lowest lying conformers corresponding to a particular secondary structure are typically within 1-2 kcal mol<sup>-1</sup> (Fig. 7). At room temperature, they would correspond to populations not differing more than by one order of magnitude. These subtle equilibria can be easily overruled by strong intramolecular forces accompanying the "collapse" of the protein into the folded structures (as mentioned above, we have recently reported that strain energies within the folded protein structures can be, exceptionally, as high as 5 kcal mol<sup>-1</sup> per amino acid residue).22 Thus, the conformers seen at experimental temperatures for the isolated tripeptides might not always be relevant for the behavior of the triplets in proteins. An example studied here is the KAM triplet/tripeptide that has a propensity for the PPII helix as an isolated tripeptide, while adopting  $\alpha$ -helical conformation in  $\sim$ 79% of its occurrence in proteins. DIC, with most  $\alpha$ -helical propensity from all studied tripeptides has 48%/28% a-helix/extended populations in proteins.

Our results show that certain peptide multiplets, as short as tripeptides, exhibit the same propensities for the specific

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secondary structure in solution in which they are preferentially found in proteins (most pronounced for pro- $\beta$ -sheet IYI and VIV). We hypothesize that these short peptides can be considered "seeds" that are important during protein folding. This compares well with our work on the WW domain<sup>7</sup> showing that low-strain parts of the WW domain(s) are the initial folding seeds despite the fact that they are not the ones most conserved within the WW protein family. Like the spark at the beginning of fire, tripeptides with an inherent secondary structure propensity could be the initiators or early-stage 'catalysts' of the folding process.

## 5. Conclusions

The experimental, bioinformatics, and computational data presented herein show that certain tripeptides have an inherent preference for certain types of secondary structure. This statement can be deconvoluted from the complex experimental and computational background characterizing their conformational behavior. This has been indicated by VCD, ECD, and NMR spectroscopies and fully supported by the quantum chemical calculations. The theory provided an unambiguous structure/ energy mapping to couple the computed data with NMR spectra and theoretically predicted VCD spectra to connect lowenergy conformers to the VCD experimental data. Some of the studied tripeptides (notably DIC( $\alpha$ ), VIV( $\beta$ ), and IYI( $\beta$ )) could be considered "folding seeds", initiating the complex and multidimensional process of protein folding. Somewhat surprisingly, only in some cases, the preference of a standalone tripeptide was the same as its behavior in proteins. This, again, suggests that the final conformation of a peptide fragment within a (folded) protein is an interplay of multiple subtle factors. In contrast, the reference CATWEAMEKCK undecapeptide has been unambiguously shown, by NMR, VCD, and ECD, to form a stable  $\alpha$ -helix in solution. A less optimistic view of the presented results may lead to the statement that the secondary structure starts to appear somewhere between 3 and 11 amino acid long peptide sequences.

## Data availability

The primary computational data as well as additional experimental data were deposited in the ESI.†

## Author contributions

M. Culka and L. Rulíšek conceived the idea for this study, and carried out initial calculations. T. Kalvoda carried out all quantum chemical calculations presented in the work and compiled, analyzed, and correlated all theoretical and experimental data. Z. Osifová and M. Dračínský carried out and interpreted NMR measurements with respect to other experimental and theoretical data. V. Andrushchenko and L. Bednárová carried out VCD and ECD experiments, respectively, and interpreted the data. P. Bouř was involved in the discussions concerning experimental and (methodological aspects of) theoretical VCD data. J. Galgonek and J. Vondrášek provided bioinformatics support. L. Rulíšek, M. Dračínský, V. Andrushchenko and T. Kalvoda wrote major parts of the manuscript. All authors assisted with editing, analysis, and interpretation.

## Conflicts of interest

There are no conflicts to declare.

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