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Hydrogen-bonding behavior of amidines in helical structure†

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Amidines are an isostere of the amide bond and are completely unexplored in peptide secondary structure. This study marks the first investigation of the structural implications of amidines in folded helices. Amidines were found to engage in hydrogen-bonding interactions that are compatible with helical structure. The protic state of the amidine is also adaptive to local interactions, able to form stronger hydrogen bonds with proton donors or form the first example of a salt bridge along the peptide backbone to stabilize the C-terminus of the helical fold. The rationalization of this behavior was aided by our discovery that the basicity of amidines within peptide backbones can be significantly lower than previously assumed for small molecules. These findings compel investigation of amidines in peptide-drug design.

1. Introduction

The prevalence of post-translational modifications in Nature suggests that expanding beyond the suite of functional groups found in the 20 canonical amino acids can impart specialized utility to peptide-based biomolecules.¹ Alteration of the peptide bond—the defining feature² of the molecule—represents a far more rarefied modification. While numerous motifs have been investigated as surrogates for the amide bond in α -helices,^{3–7} these substitutions almost always have a destabilizing effect on the structure^{8,9} except in very specific contexts.^{10–12} However, one peptide-bond isostere in particular, the amidine, has received almost no attention in peptidic molecules.^{13–15}

Amidines are distinguished from classical peptide bonds by the substitution of the carbonyl oxygen with a nitrogen atom.¹⁶ Amidines can be found within a family of natural products called ribosomally synthesized and post-translationally modified peptides (RiPPs), suggesting an evolutionary benefit may have driven the generation of biosynthetic machinery for their installation.^{17–19} Alternatively, amidines have also been proposed as primordial intermediates for the prebiotic synthesis of peptides.^{20–22} Despite these examples from Nature, studies describing the behavior of amidines in synthetic peptides are noticeably scarce.^{15,23} This unique, one-atom substitution, however, carries major implications for basicity and hydrogen-bonding behavior in the rational design of peptides (Fig. 1).^{24–27}

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The dearth of synthetic amidinopeptide examples derives from a historic lack of methods to insert amidines into peptides without side reactivity. We recently reported a synthetic approach that allowed for the first generally applicable method to site-selectively insert amidines along the peptide backbone.^{23,28}

This study represents the first report elucidating the behavior of amidines in folded protein secondary structure. This work challenges long held assumptions about the basic character of amidines and provides the first spectroscopic evidence for the malleable protic states of amidines in response to their local environment, which carries major implications for the design and drug action of vancomycin derivatives against resistant bacteria.^{13,14} Amidines are the first peptide-bond isostere to stabilize helical structure as a hydrogen-bond donor and acceptor. This single-atom substitution provides a non-covalent strategy to stabilize helices in a complimentary way to existing covalent strategies that exploit side-chain and backbone macrocyclization.^{29–31}

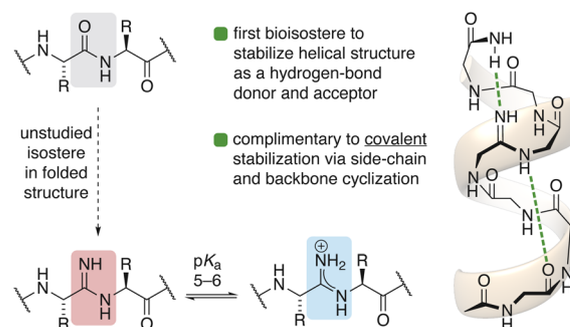


Fig. 1 Single-atom substitution with amide-like behavior.



2. Results and discussion

2.1 C-terminal effects of amidines in α -helices

To begin the investigation of how amidines might behave within canonical secondary structure, we chose to study a model α -helix first reported by Arora and coworkers (Table 1).³⁴ We were particularly interested in how the dynamic protic states of amidines might affect helicity in response to different chemistries at the C-terminus. Consider peptide **1**, which displays a C-terminal carboxylate. At pH 7.4 (phosphate buffer with 30% TFE)³⁴ **1** displays no secondary structure, as evinced by the circular dichroism (CD) spectrum of **1** that shows signatures associated with a random coil (grey-dashed line, Fig. 2). This result is not surprising given the interaction between the C-terminal carboxylate and the oxygen of the amide at Ala₇ is repulsive. We hypothesized that this interaction could be rendered attractive and stabilize the helical fold if the amide of Ala₇ were replaced with a primary amidine that, upon protonation, could form the first example of a stabilizing, backbone-to-backbone salt bridge. Accordingly, analysis of **2**, a peptide that supplies an amidine suitably placed to complement the negatively-charged carboxylate with a positively-charged amidinium, revealed a more structured peptide than **1**, with helical character indicated by negative absorbance at 222 nm in the CD spectrum (blue line, Fig. 2).

Alternatively, when the C-terminal chemistry of the initial model peptide was changed to a primary amide, **3**, the peptide maintained some helical character, presumably because an intra-strand hydrogen bond stabilizes the helical fold at the C-terminus (dotted line, Fig. 2). Remarkably, we observed that the helicity of the peptide was even further stabilized by the

presence of an amidine at Ala₇ in **4** (red line, Fig. 2). We hypothesize that this increased helical propensity was due to the stronger basicity of the amidine relative to the native amide, resulting in a stronger hydrogen bond as a result. Finally, the amidines under consideration were stable during this and all subsequent analyses (Fig. S49[†]) and do not significantly impact CD absorbance (Fig. S50[†]).

2.2 The pH-dependent behavior of amidines in peptides

An alternative explanation for the increased helicity of **4** could be rationalized in the context of structure **4-rot** (Fig. 3), in which an amidinium donates a charge-enhanced hydrogen bond to the oxygen of the C-terminal primary amide—a simple bond rotation.

To probe for this possibility, the pH of the solution was varied and the helicity of **4** was found to decrease under acidic pH and increase at more basic pH (Fig. 4A). No substantive changes in the helicity of the native all-oxoamide control peptide **3** were observed upon varying the pH over the same range (Fig. 4B), confirming that changes in helicity were likely due to the influence of the amidine moiety in **4** and not due to side-chain interactions. This behavior supports the notion of a free-base amidine, whereupon protonation of the amidine at acidic pH would disrupt the stabilizing C-terminal interaction in **4**.¹⁵ Alternatively, basic pH would promote the free-base form of the amidine, stabilizing the helical fold in **4**. These results, however, do not align with the interaction described in **4-rot** (Fig. 3).

The results above, and the proposed hydrogen-bonding interaction, led us to hypothesize that the amidine could be moved to a more internal residue within the helix without

Table 1 Sequence and circular dichroism data for controls and amidinopeptides

Peptide	Sequence	Solvent ^a	$[\theta]_{222}$	$[\theta]_{208}$	% Helicity
1 ^b	Ac-QVA RQLA  E I Y-OH	PBS	-1765	-3016	na
		30% TFE	-610	-1145	na
2	Ac-QVA RQLA  E I Y-OH	PBS	-1944	-5686	8
		10% TFE	-2153	-6960	9
		20% TFE	-7364	-10174	31
		30% TFE	-9735	-12773	42
3	Ac-QVA RQLA  E I Y-NH2	PBS	-4261	-7765	18
		10% TFE	-1544	-4998	17
		20% TFE	-7415	-10082	31
		30% TFE	-9113	-11738	38
		PBS	-2166	-5687	9
4	Ac-QVA RQLA  E I Y-NH2	10% TFE	-4740	-8972	20
		20% TFE	-8435	-11136	36
		30% TFE	-15844	-18075	67
		PBS	-1302	-2644	5
		10% TFE	-3338	-4016	14
5	Ac-QVA  RQLA E I Y-NH2	20% TFE	-7559	-8646	32
		30% TFE	-12624	-14677	54

^a 7.4 pH PBS buffer with increasing %v/v of TFE. ^b **1** remained unstructured even with the addition of TFE. See ESI, Fig. S55, for traces of all peptides at all concentrations of TFE. All peptide concentrations were standardized to the absorbance of the tyrosine residue.^{32,33} Error in measurement was 5%. The percent helicity was determined by the ratio of $[\theta]_{222}/[\theta]_{\max}$ at 20 °C. The $[\theta]_{\max}$ was determined using -23400 at a temperature of 20 °C, see eqn (2) in the ESI.



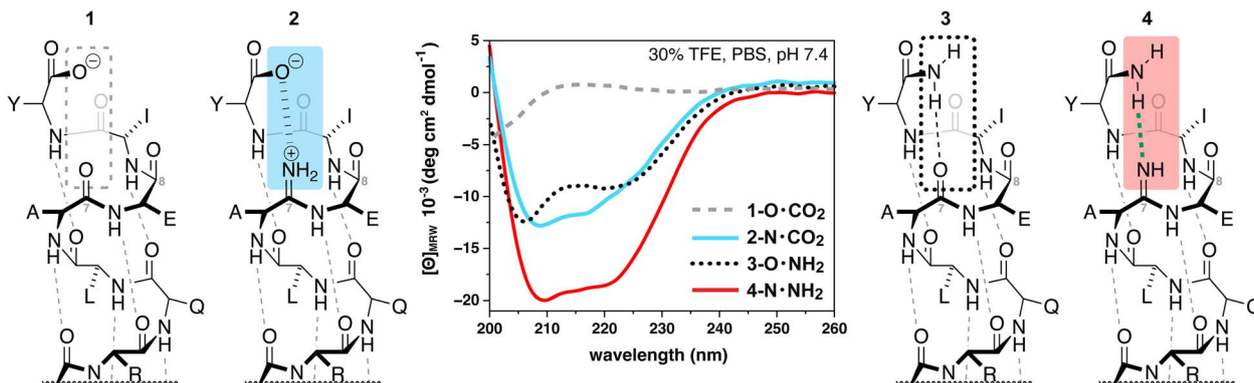


Fig. 2 Helical propensities of model peptide Ac-QVARQLAEIY-X displaying different C-terminal chemistries and interactions with the backbone, where X = OH for 1 and 2 and X = NH₂ for 3 and 4. The amidine in 1 and 4 was placed between Ala₇ and Glu₈.

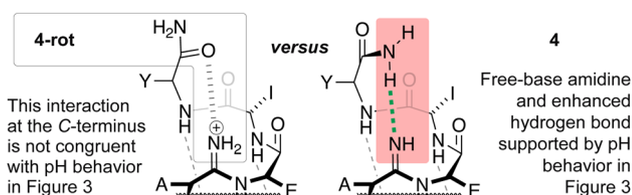


Fig. 3 Potential C-terminal interactions leading to helix stabilization. The interaction described in 4 is more consistent with the pH behavior in Fig. 4.

significant compromise to the helical fold. Accordingly, when the amidine was placed between Ala₃ and Arg₄ in the sequence, we observed a prominent helical signature in the CD spectrum for peptide 5 (Fig. 4C). Notably, the internal amidine in peptide 5 led to greater helicity than the native sequence in 3 (Fig. 4C *versus* 4B). The behavior of 5 in response to changing pH mirrors that of 4. Lower pH led to a modest decrease in helicity as determined by CD, while increased pH presumably favored the free-base of the amidine, promoting a stronger hydrogen-bonding interaction and greater observed helicity (Fig. 4C). Differences in the pH-dependent behavior between 4 and 5 likely arise due to the amidine in 4 being placed in the more disordered and solvent-exposed C-terminus compared to the more internal position in the helix for 5.

The combined results of both peptides 2 and 4 reveal that amidines are an amide-bond isostere that is tolerated in folded α -helical secondary structure and, moreover, can confer greater helical character due to a stronger hydrogen bonding interaction conferred by the more basic amidine.

2.3 Basicity and protic state of amidines in peptides

2.3.1 Basicity of amidines in peptides. The pH-dependent behavior of amidines described above and corresponding proposal of the ‘free-base’ protic state of 4 led us to reconsider some broad assumptions about amidine basicity. The amidine is generally considered to be highly basic ($pK_a \approx 12$), where these charged groups are associated with poor pharmacokinetic properties in small molecules.^{26,35,36} These assertions appear to be supported by the reported pK_a values for benzamidine and drug molecules in which the amidine decorates the periphery of the core.^{26,37} Thus, amidines have traditionally been studied in settings in which solvent exposure promotes a protonated state. In contrast, an amidine within a peptide backbone would be significantly more buried and solvated to a lesser degree. Accordingly, we measured a pK_a between 5 and 6 when an amidine was placed within an Ac-Phe-Ala-NH₂ dipeptide (6, Fig. 5). It should be noted that opportunities for intramolecular hydrogen bonding of the amidine in 6 with the amides that

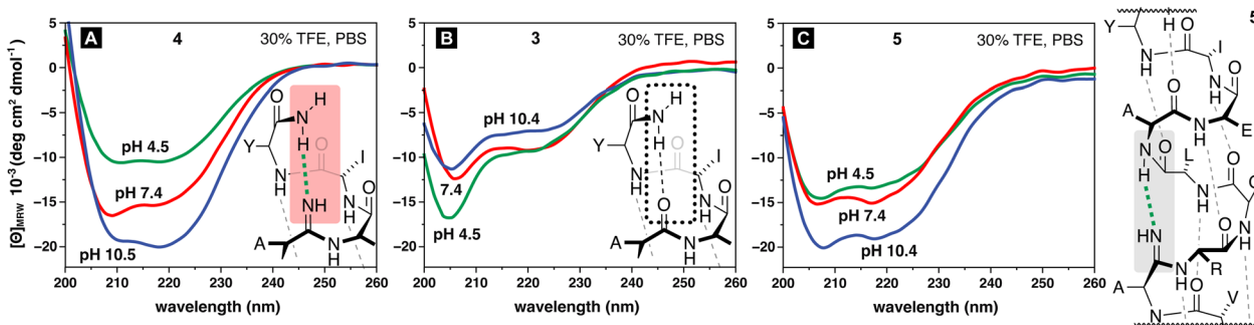


Fig. 4 pH-dependent behavior and helical propensities of model peptides based on Ac-QVARQLAEIY-NH₂. CD curves measured at 20 °C for (A) peptide 4 (B) peptide 3, and (C) peptide 5 in PBS buffer at pH 4.5, 7.4, and 10.5 with the addition of 30% TFE as a co-solvent. The amidine in 4 and 3 was placed between Ala₇ and Glu₈. The amidine in 5 was placed between Ala₃ and Arg₄.



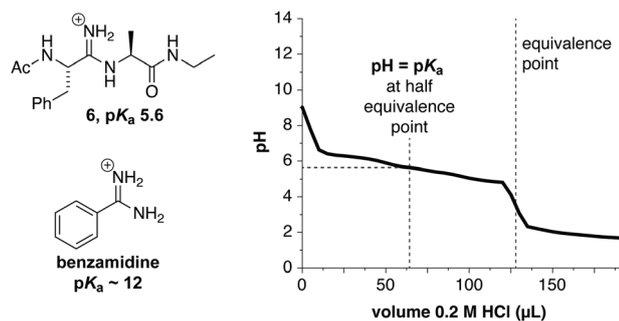


Fig. 5 Titration experiment to estimate the pK_a of an amidine in peptide structure and comparison to a more solvent-exposed benzamidine.

bracket it can lead to lower measured pK_a values.³⁸ Thus, dipeptide **6** should be regarded as a lower limit of the pK_a in folded helical structure.

The potential for more variable basic properties for amidines is in stark contrast to the related guanidine functional group in the side-chain of arginine, which has been spectroscopically demonstrated to be invariably protonated and charged ($pK_a > 12$) in proteins, even when buried within hydrophobic micro-environments.³⁹ This new understanding of amidine pK_a behavior, however, lends additional support to the work of Boger and coworkers concerning the adaptive protic states of amidines in the function of so-called maxamycins against antibiotic resistant Gram-positive bacteria.^{40,41}

2.3.2 NMR characterization of amidine protic state. To further characterize the protic state of the amidine by NMR, we next exploited our synthetic approach²⁸ for facile installation of ^{15}N (from relatively inexpensive $^{15}\text{NH}_4\text{OAc}$) at the ^2N -imino position of the amidine. This is a notable advantage of amidines compared to the amide or traditional ester and thioamide isosteres. Facile installation of ^{15}N at the ^2N -imino position of the amidine negates the need for costly ^{15}N enriched amino acids to introduce this probe nucleus into amides and thioamides. Further, the amidine furnishes a spin 1/2 ^{15}N nucleus to probe local structure and coordination in solution-phase experiments, in contrast to oxygen and sulfur nuclei which are both quadrupolar and only amenable to solid-state NMR experiments.^{42,43} Moreover, the prominent signal from the ^{15}N spin label confirms the installation of the amidine along the peptide backbone and provides further confidence in the success of the synthesis.

Spectroscopic evidence of the protic states of the amidine in all three peptides—**2**, **4**, and **5**—was gathered using ^{15}N - ^1H HSQC correlation spectroscopy (Fig. 6). The chemical shift of the enriched ^{15}N in the amidine was easily identifiable as a result of its signal intensity relative to the other amides at natural abundance.

To limit effects due to fast exchange of protons, initial experiments were conducted at pH 4.5, where all peptides were shown to have helical character (by CD (Fig. 4) and by NMR (Fig. 7 *vide infra*)). For the C-terminal amidines in **2** and **4**, we observed two cross peaks for the ^{15}N label at 109 ppm,

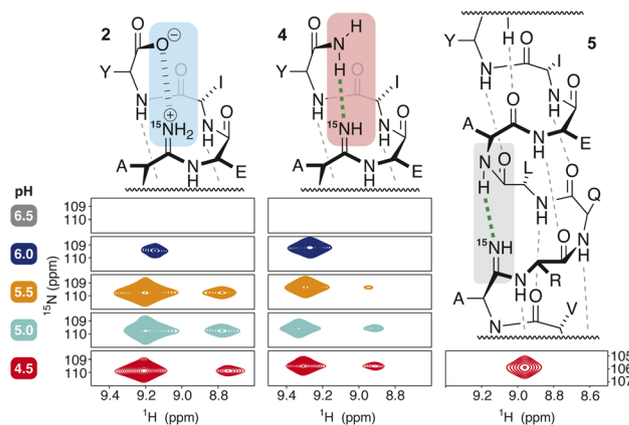


Fig. 6 NMR pH titration of model peptides at 20 °C in PBS buffer with the addition of 30% TFE as a co-solvent.

indicating that two ^1H were bound to the ^2N -imino nitrogen of the amidine at pH 4.5. This NH_2^+ protonation state supports the notion of an intra-strand salt bridge proposed for **2**. Given the pK_a value of 5–6 we estimated for an amidine in a peptide backbone (Fig. 5), the observation of a doubly protonated amidine in **4** at pH 4.5 was not unexpected. These observations, however, are in marked contrast to the results for **5** (Fig. 6), which displayed only one crosspeak, indicating the unprotonated, free-base form. We hypothesize that the double protonation of the amidine in **4** arises from its position in the more disordered C-terminus of the peptide (Fig. 7), leading to greater solvent exposure, providing more opportunity for hydrogen-bonding interactions with protons from solvent. The amidine in **5**, however, is in the middle of the helix and appears to be protected from solvation.

We next attempted to raise the pH to see if we could observe changes in the protic state for **2** and **4** (Fig. 6). Unfortunately, fast-exchange processes lead to loss of the ^{15}N - ^1H cross peaks within only 2 pH units, which limited the scope of the analysis. This observation is not uncommon in protein NMR, and lowering the temperature of the analysis to 5 °C did not restore the signal. Several interesting observations still provided insight into the structure however. In the case of **2**, the ratio of the two cross peaks remains constant as the pH is increased, until the signals are lost to fast exchange at pH 6.0. We propose that this observation supports the presence of an NH_2^+ moiety that persists as pH increases, consistent with our proposed intra-strand salt bridge **2**. Conversely, for **4** the cross peak for one of the protons on the ^{15}N loses intensity as the pH increases, accompanied by a shift in the remaining proton, until complete loss of signal at pH 6.5. We propose that this observation supports the notion of a mono-protonated NH moiety that becomes protonated to a greater extent at lower pH, consistent with our proposed free-base amidine in **4**.

2.4 NMR characterization of helical structure

This study is the first to characterize the behavior of amidines in folded peptide structure. Thus, to both further interrogate the results of the CD experiments and identify the effects of



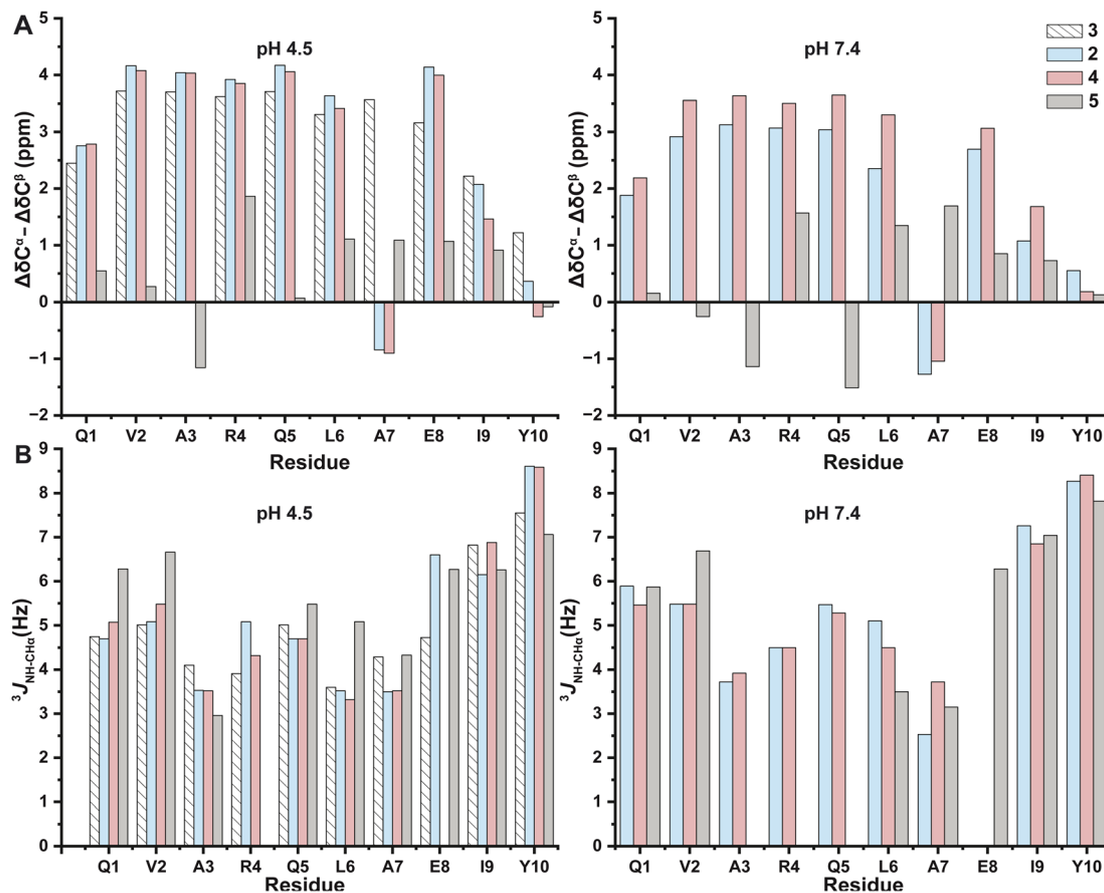


Fig. 7 Secondary structure analysis for 2, 3, 4, and 5 using (A) the difference between experimentally observed $^{13}\text{C}^\alpha$ and $^{13}\text{C}^\beta$ chemical shifts and reference shifts for a random coil at pH 4.5 (left) and pH 7.4 (right) and (B) the $^3J_{\text{NH-CH}^\alpha}$ coupling constants at pH 4.5 (left) and pH 7.4 (right).^{44,45}

amidine incorporation on NMR secondary-structure analysis, ^{13}C - ^1H and ^{15}N - ^1H HQSCs, NOESY, and TOCSY experiments were used to fully assign the chemical shifts of 2, 4, and 5 (see Tables S1 and S2 in ESI† for residue assignments).

Analysis of the high-field (800 MHz) 2D-NOESY NMR data was used to confirm peptide helicity. NOE correlations along the peptide backbone are key signatures of helical structure (Table 2). Specifically, sequential (NN) and ${}_2\text{N}$ ($i, i + 1$) and non-sequential ${}_3\text{N}$ ($i, i + 3$) cross-peaks are indisputable evidence for helical structure. Although spectral overlap inhibits the observation of all cross-peaks, a significant number of signature cross-peaks were observed for the peptides under investigation, further validating helical structure. NOE correlation maps and an NOE correlation table are provided in ESI (Fig. S36 and S39†).

Following the chemical shift assignments, we characterized the residue-specific helicity within each peptide by two independent ways. The first method used the differences between the experimentally observed $^{13}\text{C}^\alpha$ and $^{13}\text{C}^\beta$ chemical shifts and the corresponding chemical shifts for a random coil.^{46,47} $^{13}\text{C}^\alpha$ and $^{13}\text{C}^\beta$ chemical shifts have been shown to be extremely sensitive and accurate indicators of secondary structure, even more accurate than H^α chemical shifts when distinguishing an α -helix from a random coil.^{46–48} When comparing experimental chemical shifts against corresponding random coil chemical

shifts, positive $\Delta\delta\text{C}^\alpha - \Delta\delta\text{C}^\beta$ values are indicative of α -helical structure (and negative $\Delta\delta\text{C}^\alpha - \Delta\delta\text{C}^\beta$ values indicate β -sheet structure).^{49–51} Thus, based on this method of analysis, both 2 and 4 display helical structure along the sequence with a notable exception for Ala₇, which is the residue that bears the amidine for both peptides (Fig. 7A). The distinctly negative $\Delta\delta\text{C}^\alpha - \Delta\delta\text{C}^\beta$ value for Ala₇ in these peptides is not reliable for secondary structure analysis, however, because the corresponding random-coil chemical shifts are referenced for native oxoamide amino acids. Thus, the chemical shift of an Ala₇ amidine-containing amino acid is therefore not suitable for the analysis *via* these methods.^{52,53}

The aberrant $\Delta\delta\text{C}^\alpha - \Delta\delta\text{C}^\beta$ chemical shift values imposed by the amidine functional group have substantially greater impact on the analysis of 5, which bears the internal amidine at Ala₃ (Fig. 7). The magnitude of the $\Delta\delta\text{C}^\alpha - \Delta\delta\text{C}^\beta$ values are generally lower relative to 2 and 4 across the sequence and highly variable around the amidine itself. The negative $\Delta\delta\text{C}^\alpha - \Delta\delta\text{C}^\beta$ for Gln₇ is particularly noteworthy and not congruent with the CD data described above. Because the $\Delta\delta\text{C}^\alpha - \Delta\delta\text{C}^\beta$ analysis is based on empirically derived chemical shifts for α -helices, it is unsurprising that these results are not appropriate to a novel and non-standard backbone modification such as the amidine. We therefore sought to interrogate the structure using a different



Table 2 Observable NOE correlation cross-peaks along the backbone for sequential NN ($i, i + 1$) and d_2N ($i, i + 1$) crosspeaks. In addition to the non-sequential medium range NOEs, d_2N ($i, i + 3$) and d_2N ($i, i + 4$) to support α -helical conformation for amidinopeptides at pH 4.5 and 7.4

Peptide	pH 4.5				pH 7.4				
	NN	d_2N ($i, i + 1$)	d_2N ($i, i + 3$)	d_2N ($i, i + 4$)	NN	d_2N ($i, i + 1$)	d_2N ($i, i + 3$)	d_2N ($i, i + 4$)	
2	Q1 \rightarrow V2	Q1 \rightarrow V2	V2 \rightarrow Q5	V2 \rightarrow L6	Q1 \rightarrow V2	Q1 \rightarrow V2	V2 \rightarrow Q5		
	V2 \rightarrow A3	V2 \rightarrow A3	A3 \rightarrow L6		V2 \rightarrow A3	V2 \rightarrow A3	A3 \rightarrow L6		
	R4 \rightarrow Q5	Q5 \rightarrow L6	R4 \rightarrow A7		R4 \rightarrow Q5	Q5 \rightarrow L6	R4 \rightarrow A7		
	Q5 \rightarrow L6	L6 \rightarrow A7	L6 \rightarrow I9		L6 \rightarrow A7	L6 \rightarrow A7			
	L6 \rightarrow A7	E8 \rightarrow I9	A7 \rightarrow Y10		I9 \rightarrow Y10	E8 \rightarrow I9			
	A7 \rightarrow E8	I9 \rightarrow Y10				I9 \rightarrow Y10			
	E8 \rightarrow I9								
	I9 \rightarrow Y10								
	4	Q1 \rightarrow V2	Q1 \rightarrow V2	V2 \rightarrow Q5	V2 \rightarrow L6	Q1 \rightarrow V2	Q1 \rightarrow V2	Q1 \rightarrow R4	V2 \rightarrow L6
		V2 \rightarrow A3	R4 \rightarrow Q5	A3 \rightarrow L6		V2 \rightarrow A3	V2 \rightarrow A3	V2 \rightarrow Q5	
R4 \rightarrow Q5		Q5 \rightarrow L6	R4 \rightarrow A7		R4 \rightarrow Q5	L6 \rightarrow A7	R4 \rightarrow A7		
Q5 \rightarrow L6		L6 \rightarrow A7	Q5 \rightarrow E8		Q5 \rightarrow L6	I9 \rightarrow Y10	L6 \rightarrow I9		
L6 \rightarrow A7		E8 \rightarrow I9	L6 \rightarrow I9		L6 \rightarrow A7				
A7 \rightarrow E8		I9 \rightarrow Y10			I9 \rightarrow Y10				
E8 \rightarrow I9									
I9 \rightarrow Y10									
5		Q1 \rightarrow V2	Q1 \rightarrow V2			L6 \rightarrow A7	Q1 \rightarrow V2	A7 \rightarrow Y10	
		V2 \rightarrow A3	V2 \rightarrow A3			A7 \rightarrow E8	E8 \rightarrow I9		
	Q5 \rightarrow L6	Q5 \rightarrow L6			E8 \rightarrow I9	I9 \rightarrow Y10			
	L6 \rightarrow A7	L6 \rightarrow A7			I9 \rightarrow Y10				
	E8 \rightarrow I9	E8 \rightarrow I9							

spectroscopic signature that was not dependent on chemical shift.

The ambiguities with respect to differences in the chemical shift displayed by the amidine compared to the native amide drove us to pursue a second method to characterize the secondary structure. Specifically, the dihedral angle between the amide NH and the CH^α (the ϕ angle) is another signature of helical structure and is independent of chemical shift. In a freely rotating, random coil, the $^3J_{HN-CH^\alpha}$ coupling for the ϕ -angle is >6 Hz. The ϕ -angle in an α -helix, however, is an acute -57° , which corresponds to a $^3J_{HN-CH^\alpha}$ coupling value of 3–6 Hz (the $^3J_{HN-CH^\alpha}$ coupling value in a β -sheet conformation is >8 Hz).⁵⁴

Thus, the residue-specific $^3J_{HN-CH^\alpha}$ values for each peptide 2, 4, and 5 indicated helical structure along the sequence (Fig. 7B and Table S3†). Residues near the N- and C-termini are likely to be the most disordered, as indicated by $^3J_{HN-CH^\alpha}$ values that are >6 Hz. Importantly, Ala₇ in both 2 and 4 displayed helical character, in agreement with the CD experiments (Fig. 2), and confirming our suspicions that the negative $\Delta\delta C^\alpha - \Delta\delta C^\beta$ values with respect to Ala₇ in Fig. 7A were an artifact of the unique chemical shifts of the amidine in that particular difference analysis.

The $^3J_{HN-CH^\alpha}$ coupling value for Ala₃ in 5 similarly indicates a helical conformation. Residues on the N-terminal side of Ala₃ (Gln₁ and Val₂) are more disordered than the analogous residues in 2 and 4. Residues on the C-terminal side of Ala₃ in 5 (Arg₄ and Gln₅) are subject to fast-exchange process that broaden these signals below the limit of detection and prevent analysis. These exchange processes are known to increase in

severity as the pH of the sample approaches 7. Accordingly, we observe loss of signal for Ala₃, Arg₄ and Gln₅ at pH 7.4 (Fig. 7B). These exchange processes may indicate that the peptide is sampling different conformations, perhaps exploring an $i + 4 \rightarrow i$ hydrogen bond between Leu₆ and Ala₃ (α -helical) and an $i + 3 \rightarrow i$ hydrogen bond between Gln₅ and Ala₃ (3_{10} helix). This conformational change would impact the intervening Arg₄ and Gln₅ most significantly.

3. Conclusion

This work is the first study to investigate the structural implications of amidines in helical peptide structure. Amidines are an historically ignored amide-bond isostere, but recently reported methods for installation into peptide backbones^{23,28,55} now make it possible to investigate these groups. Collectively, the results of this work indicate that amidine isosteres are not only tolerated in folded helical structure, but make interactions that are intuitive and amenable to design. These findings compel investigation of amidines in peptide-drug design, particularly in light of the fact that the pK_a of amidines in peptides may be lower than previously assumed, thereby improving the potential pharmacokinetic profile.

More specifically, helicity is most tolerant of amidines at the C-terminus, likely due to the greater flexibility associated with the fraying ends of the helix. Depending on the C-terminal chemistry, the amidine can adapt its protic state to its local environment. In the case of a C-terminal carboxylate, an amidinium forms the first example of an intrachain salt bridge. Alternatively, in the case of a C-terminal hydrogen-bond donor,



a free-base amidine can form a stronger hydrogen bond than a traditional amide to stabilize the helical fold. The amidine can also be inserted in the middle of the peptide without detrimental loss of overall helical character, although the residues around the amidine may experience greater disorder.

Evidence for the nature of the protic state of the amidine was greatly aided by the adventitious spin $1/2$ ^{15}N nucleus that could be readily inserted during peptide synthesis. The ability to probe local interactions around a particular backbone site using solution-phase NMR techniques makes amidines a powerful structural tool.

4. Experimental

All experimental details and characterization data are provided free of charge in the electronic ESI.†

Data availability

Data for this article, including synthetic and experimental procedures, CD and NMR spectroscopic data for characterization of conformation and structure, and other supporting experiments has been included in the ESI† and available free of charge on the publisher's website for this article.

Author contributions

B. V., E. A. O. and B. J. W. conceived of the work. E. A. O collected and analyzed CD data. J. A. P. collected and analyzed NMR data with E. A. O. All authors contributed to the writing and proofing of the final manuscript.

Conflicts of interest

There are no conflicts to declare.

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References

- H. Lin and K. S. Carroll, *Chem. Rev.*, 2018, **118**, 887–888.
- E. Fischer and B. Dtsch, *Chem. Ges.*, 1906, **39**, 530–610.
- A. Choudhary and R. T. Raines, *ChemBioChem*, 2011, **12**, 1801–1807.
- S. Kumari, A. V. Carmona, A. K. Tiwari and P. C. Trippier, *J. Med. Chem.*, 2020, **63**, 12290–12358.
- N. A. Meanwell, *J. Med. Chem.*, 2011, **54**, 2529–2591.
- N. A. Meanwell, *J. Agric. Food Chem.*, 2023, **71**, 18087–18122.
- J. J. Jackson, A. C. Siegmund, W.-J. Bai, A. B. Reed, A. B. Birkholz, I. D. G. Campuzano, A. Créquer-Grandhomme, R. Hu, R. V. Modak, A. Sudom, N. Javier, C. Sanders, M.-C. Lo, F. Xie, V. J. Cee, P. Manzanillo and J. G. Allen, *J. Med. Chem.*, 2023, **66**, 16120–16140.
- J. Byerly-Duke and B. VanVeller, *Org. Lett.*, 2024, **26**, 1452–1457.
- A. Reiner, D. Wildemann, G. Fischer and T. Kiefhaber, *J. Am. Chem. Soc.*, 2008, **130**, 8079–8084.
- J. H. Miwa, L. Pallivathucal, S. Gowda and K. E. Lee, *Org. Lett.*, 2002, **4**, 4655–4657.
- C. R. Walters, D. M. Szantai-Kis, Y. Zhang, Z. E. Reinert, W. S. Horne, D. M. Chenoweth and E. J. Petersson, *Chem. Sci.*, 2017, **8**, 2868–2877.
- C. Wu, H. N. Hoang, T. A. Hill, J. Lim, W. M. Kok, K. Akondi, L. Liu and D. P. Fairlie, *Chem. Commun.*, 2022, **58**, 12475–12478.
- A. Okano, R. C. James, J. G. Pierce, J. Xie and D. L. Boger, *J. Am. Chem. Soc.*, 2012, **134**, 8790–8793.
- J. Xie, A. Okano, J. G. Pierce, R. C. James, S. Stamm, C. M. Crane and D. L. Boger, *J. Am. Chem. Soc.*, 2012, **134**, 1284–1297.
- S. Huh, S. D. Appavoo and A. K. Yudin, *Org. Lett.*, 2020, **22**, 9210–9214.
- D. L. Boger, *J. Org. Chem.*, 2017, **82**, 11961–11980.
- W. J. K. Crone, N. M. Vior, J. Santos-Aberturas, L. G. Schmitz, F. J. Leeper and A. W. Truman, *Angew. Chem., Int. Ed.*, 2016, **55**, 9639–9643.
- K. A. Clark and M. R. Seyedsayamdost, *J. Am. Chem. Soc.*, 2022, **144**, 17876–17888.
- A. H. Russell, N. M. Vior, E. S. Hems, R. Lactret and A. W. Truman, *Chem. Sci.*, 2021, **12**, 11769–11778.
- C. S. Foden, S. Islam, C. Fernández-García, L. Maugeri, T. D. Sheppard and M. W. Powner, *Science*, 2020, **370**, 865–869.
- J. Fairchild, S. Islam, J. Singh, D.-K. Bučar and M. W. Powner, *Science*, 2024, **383**, 911–918.
- J. Singh, D. Whitaker, B. Thoma, S. Islam, C. S. Foden, A. E. Aliev, T. D. Sheppard and M. W. Powner, *J. Am. Chem. Soc.*, 2022, **144**, 10151–10155.
- B. J. Wall, K. K. Sharma, E. A. O'Brien, A. Donovan and B. VanVeller, *J. Am. Chem. Soc.*, 2024, **146**, 11648–11656.
- A. A. Aly, S. Bräse and M. A.-M. Gomaa, *Arkivoc*, 2018, **6**, 85–138.
- A. A. Aly and A. M. Nour-El-Din, *Arkivoc*, 2008, **1**, 153–194.
- P. Y. S. Lam, C. G. Clark, R. Li, D. J. P. Pinto, M. J. Orwat, R. A. Galemno, J. M. Fevig, C. A. Teleha, R. S. Alexander, A. M. Smallwood, K. A. Rossi, M. R. Wright, S. A. Bai, K. He, J. M. Luetgen, P. C. Wong, R. M. Knabb and R. R. Wexler, *J. Med. Chem.*, 2003, **46**, 4405–4418.
- J. Oszczapowicz and J. Jaroszevska-Manaj, *J. Chem. Soc., Perkin Trans. 2*, 1991, 1677–1680.



- 28 E. A. O'Brien, K. K. Sharma, J. Byerly-Duke, L. A. Camacho III and B. VanVeller, *J. Am. Chem. Soc.*, 2022, **144**, 22397–22402.
- 29 T. A. Hill, N. E. Shepherd, F. Diness and D. P. Fairlie, *Angew. Chem., Int. Ed.*, 2014, **53**, 13020–13041.
- 30 D. A. Guarracino, J. A. Riordan, G. M. Barreto, A. L. Oldfield, C. M. Kouba and D. Agrinoni, *Chem. Rev.*, 2019, **119**, 9915–9949.
- 31 S. E. Miller, A. M. Watkins, N. R. Kallenbach and P. S. Arora, *Proc. Natl. Acad. Sci. U.S.A.*, 2014, **111**, 6636–6641.
- 32 E. Gasteiger, C. Hoogland, A. Gattiker, S. Duvaud, M. R. Wilkins, R. D. Appel and A. Bairoch, in *Protein Identification and Analysis Tools on the ExPASy Server*, ed. J. M. Walker, Humana Press, 2005, pp. 571–607.
- 33 N. J. Anthis and G. M. Clore, *Protein. Sci.*, 2013, **22**, 851–858.
- 34 D. Wang, K. Chen, J. L. Kulp and P. S. Arora, *J. Am. Chem. Soc.*, 2006, **128**, 9248–9256.
- 35 T. M. Kamenecka, Y.-J. Park, L. S. Lin, S. de Laszlo, E. D. McCauley, G. Van Riper, L. Egger, U. Kidambi, R. A. Mumford, S. Tong, W. Tang, A. Colletti, Y. Teffera, R. Stearns, M. MacCoss, J. A. Schmidt and W. K. Hagmann, *Bioorg. Med. Chem. Lett.*, 2004, **14**, 2323–2326.
- 36 Y. M. Choi-Sledeski and C. G. Wermuth, *The Practice of Medicinal Chemistry*, Academic Press, San Diego, 4th edn, 2015, pp. 657–696.
- 37 J. V. Greenhill and P. Lue, *Prog. Med. Chem.*, 1993, **30**, 203–326.
- 38 D. Kanamori, A. Furukawa, T.-a. Okamura, H. Yamamoto and N. Ueyama, *Org. Biomol. Chem.*, 2005, **3**, 1453–1459.
- 39 C. A. Fitch, G. Platzer, M. Okon, B. E. Garcia-Moreno and L. P. McIntosh, *Protein Sci.*, 2015, **24**, 752–761.
- 40 Z.-C. Wu and D. L. Boger, *Acc. Chem. Res.*, 2020, **53**, 2587–2599.
- 41 M. J. Moore, P. Qin, D. J. Keith, Z.-C. Wu, S. Jung, S. Chatterjee, C. Tan, S. Qu, Y. Cai, R. L. Stanfield and D. L. Boger, *J. Am. Chem. Soc.*, 2023, **145**, 12837–12852.
- 42 S. L. Carnahan, B. J. Lampkin, P. Naik, M. P. Hanrahan, I. I. Slowing, B. VanVeller, G. Wu and A. J. Rossini, *J. Am. Chem. Soc.*, 2019, **141**, 441–450.
- 43 A. V. Wijesekara, A. Venkatesh, B. J. Lampkin, B. VanVeller, J. W. Lubach, K. Nagapudi, I. Hung, P. L. Gor'kov, Z. Gan and A. J. Rossini, *Chem.–Eur. J.*, 2020, **26**, 7881–7888.
- 44 G. Merutka, H. Jane Dyson and P. E. Wright, *J. Biomol. NMR*, 1995, **5**, 14–24.
- 45 M. Kjaergaard and F. M. Poulsen, *J. Biomol. NMR*, 2011, **50**, 157–165.
- 46 D. S. Wishart and B. D. Sykes, *J. Biomol. NMR*, 1994, **4**, 171–180.
- 47 Y. Wang and O. Jardetzky, *Protein Sci.*, 2002, **11**, 852–861.
- 48 D. Eliezer, in *Characterizing Residual Structure in Disordered Protein States Using Nuclear Magnetic Resonance*, ed. Y. Bai and R. Nussinov, Humana Press, Totowa, NJ, 2006, pp. 49–67.
- 49 D. Eliezer, E. Kutluay, R. Bussell Jr and G. Browne, *J. Mol. Biol.*, 2001, **307**, 1061–1073.
- 50 S. Spera and A. Bax, *J. Am. Chem. Soc.*, 1991, **113**, 5490–5492.
- 51 D. S. Wishart, B. D. Sykes and F. M. Richards, *J. Mol. Biol.*, 1991, **222**, 311–333.
- 52 A. Bundi and K. Wüthrich, *Biopolymers*, 1979, **18**, 285–297.
- 53 D. G. Isom, C. A. Castañeda, B. R. Cannon and B. García-Moreno E, *Proc. Natl. Acad. Sci. U. S. A.*, 2011, **108**, 5260–5265.
- 54 L. Mander and H.-W. Liu, *Comprehensive Natural Products II: Chemistry and Biology*, Elsevier, 2010, vol. 1.
- 55 S. Liu, Y. Tang, S. Chen, X. Li and H. Liu, *Angew. Chem., Int. Ed.*, 2024, **63**, e202407952.

